

# HRAS as a potential therapeutic target of salirasib RAS inhibitor in bladder cancer

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**Abstract.** The active form of the small GTPase RAS binds to downstream effectors to promote cell growth and proliferation. RAS signal enhancement contributes to tumorigenesis, invasion, and metastasis in various different cancers. HRAS proto-oncogene GTPase (HRAS), one of the RAS isoforms, was the first human oncogene for which mutations were reported in T24 bladder cancer (BC) cells in 1982, and HRAS mutation or upregulation has been reported in several cancers. According to data from The Cancer Genome Atlas, HRAS expression was significantly upregulated in clinical BC samples compared to healthy samples ( $P=0.0024$ ). HRAS expression was also significantly upregulated in BC with HRAS mutation compared to patients without HRAS mutation ( $P<0.0001$ ). The tumor suppressive effect of salirasib, a RAS inhibitor, has been reported in several cancer types, but only at relatively high concentrations. As such, RAS inhibitors have not been used for clinical applications. The aim of the current study was to investigate the therapeutic potential of targeting HRAS using salirasib and small interfering RNA (siRNA) and to characterize the mechanism by which HRAS functions using recently developed quantitative *in vitro* proteome-assisted multiple reaction monitoring for protein absolute quantification (iMPAQT), in BC cells. iMPAQT allows measurement of the absolute abundance of any human protein with the high quantitative accuracy. Salirasib and siRNA targeting of HRAS inhibited cell proliferation, migration and invasion in HRAS wild type and HRAS-mutated cell lines. Proteomic analyses revealed that several metabolic pathways, including the oxida-

tive phosphorylation pathway and glycolysis, were significantly downregulated in salirasib-treated BC cells. However, the expression levels of hexokinase 2, phosphoglycerate kinase 1, pyruvate kinase, muscle (PKM)1, PKM2 and lactate dehydrogenase A, which are downstream of RAS and target genes of hypoxia inducible factor-1 $\alpha$ , were not notably downregulated, which may explain the high concentration of salirasib required to inhibit cell viability. These findings provide insight into the mechanisms of salirasib, and suggest the need for novel therapeutic strategies to treat cancers such as BC.

## Introduction

Bladder cancer (BC) was the 5th most commonly diagnosed cancer and the 8th most common cause of cancer-associated mortality among the 40 European Union countries in 2012. In that same year, 429,800 new cases of BC were diagnosed, and 165,000 patients succumbed to BC worldwide (1,2). The 5-year survival rate of patients with BC has improved by only a small percentage during the last 30 years according to the National Cancer Institute program Surveillance, Epidemiology and End Results (3). One factor in the lack of improvement in BC survival rates is the limited efficacy of cisplatin-based combination chemotherapy (4). Thus, innovative therapeutic strategies are required to improve BC outcomes.

RAS proteins are small molecular weight GTPases that couple extracellular signals to intracellular effector pathways. Mammalian cells encode three closely related RAS proteins, HRas proto-oncogene GTPase (HRAS), NRAS proto-oncogene GTPase (NRAS) and KRAS proto-oncogene GTPase (KRAS), which have critical roles in fundamental cellular processes, including proliferation, survival, differentiation, motility and transcription (5). The RAS pathway is one of the most commonly deregulated pathways in human cancer (6), and activating mutations in RAS genes occur in ~30% of all tumors (7). These mutations typically render RAS as constitutively GTP-bound, resulting in activation of downstream effector pathways regardless of extracellular stimulation (6). Substitution of glycine for valine at amino acid 12 (G12V) is one of the most frequently observed RAS mutations that interferes with GTPase-activating protein-mediated GTP hydrolysis, leading to excess amounts of active GTP-bound RAS. Notably, the type of mutated RAS gene (HRAS, KRAS or

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*NRAS*) varies depending on the tumor type; *KRAS* mutations are frequently detected in pancreatic carcinoma (80-90%) and colorectal carcinoma (30-60%), whereas *HRAS* mutations are frequent in BC (7-66%) and thyroid cancer (0-60%) (8).

RAS is considered 'undruggable' because the RAS protein lacks a druggable binding pocket (9,10). Additionally, development of specific and competitive nucleotide inhibitors is challenging, because RAS binds nucleotide ligands with high affinity (10). To overcome these challenges, the RAS antagonist salirasib, also termed farnesylthiosalicylate, was designed to competitively inhibit attachment of GTP-bound RAS to the plasma membrane, which in turn inactivates RAS signaling (11). Salirasib inhibits all RAS isoforms and inhibits the growth of RAS-driven cancer (11,12). Although applications of salirasib have been tested in several clinical trials for cancers other than BC (13-16), it exhibited insufficient tumor suppressive effects in trials in which salirasib was the single agent. In addition, relatively high concentrations of salirasib were required to achieve sufficient tumor suppressive effects (17,18). Therefore, a more detailed examination of the effects of salirasib is required to understand the mechanism by which salirasib acts on RAS. Recently, Matsumoto *et al.* (19) developed a targeted proteomics platform, *in vitro* proteome-assisted multiple reaction monitoring for protein absolute quantification (iMPAQT), that analyzes 18,000 human recombinant proteins to enable absolute protein quantification on a genome-wide scale (19) and overcome limitations in quantitative accuracy, reproducibility, and analysis speed associated with conventional analysis methods. Use of iMPAQT allows large-scale and accurate assessment of protein abundances that can influence cellular phenotypes.

In the current study, the therapeutic potential of *HRAS* knockdown by salirasib or RNA interference was investigated in two BC cell lines (T24 cells with *HRAS* G12V mutation and BOY cells without *HRAS* mutation). Furthermore, newly developed quantitative proteome analysis of BC cells treated with salirasib was performed to elucidate the mechanisms underlying the actions of salirasib toward *HRAS*.

## Materials and methods

**Analysis in the BC cohort of The Cancer Genome Atlas (TCGA).** Sequencing data were available for 407 BC samples and 19 normal bladder epithelial samples in TCGA database ([tcga-data.nci.nih.gov/tcga/](http://tcga-data.nci.nih.gov/tcga/)). We used TCGA to analyze *HRAS* mRNA expression levels in normal and BC tissues and to evaluate differences in *HRAS* mRNA expression levels according to *HRAS* mutational status. RNA-Seq by Expectation Maximization software was used for gene expression quantification (20). Full sequencing information, somatic mutation information, and clinical information were acquired using UCSC Xena ([xena.ucsc.edu/](http://xena.ucsc.edu/)) and TCGA. The current study meets publication guidelines provided by TCGA ([cancergenome.nih.gov/publications/publicationguidelines](http://cancergenome.nih.gov/publications/publicationguidelines)).

**Cell culture and RNA extraction.** Four human BC cell lines were used. T24, KK47 and UMUC cells, which were obtained from the American Type Culture Collection (Manassas, VA, USA), and BOY cells, which were established in our laboratory from a 66-year-old Asian male patient diagnosed with stage III BC

with lung metastasis. These cell lines were maintained in the minimum essential Eagle's medium (Sigma-Aldrich; Merck KGaA, Darmstadt, Germany), containing 10% fetal bovine serum (Equitech-Bio, Inc., Kerrville, TX, USA), 50  $\mu$ g/ml streptomycin, and 50 U/ml penicillin in a humidified atmosphere of 95% air/5% CO<sub>2</sub> at 37°C. Total RNA was isolated using Isogen (Nippon Gene Co., Ltd., Tokyo, Japan) according to the manufacturer's protocol. The integrity of the RNA was checked with an RNA 6000 Nano assay kit and a 2100 Bioanalyzer (Agilent Technologies, Inc., Santa Clara, CA, USA).

**Reverse transcription-quantitative polymerase chain reaction (RT-qPCR).** A SYBR-Green qPCR-based array approach was used for RT-qPCR. RT was performed using the TaqMan High-Capacity cDNA Reverse Transcription Kit (cat. no. 4368814; Applied Biosystems; Thermo Fisher Scientific, Inc., Waltham, MA, USA) under the incubation conditions (25°C for 10 min, 37°C for 120 min and 85°C for 5 min) according to the manufacturer's instructions. The primer set for determination of mRNA expression levels was as follows: *HRAS*, forward, 5'-ATGACGGAATATAAGCTGGTGGT-3' and reverse, 5'-GGCACGCTCCCCATCAATG-3'; hypoxia inducible factor-1 $\alpha$  (*HIF-1 $\alpha$* ), forward, 5'-GAACGTCGAAAAGAAA GTCTCG-3' and reverse, 5'-CCTTATCAAGATGCGAACTC ACA-3'; glucuronidase  $\beta$  (*GUSB*), forward, 5'-CGTCCCACC TAGAATCTGCT-3' and reverse, 5'-TTGCTCACAAAGGT CACAGG-3'. The experimental procedures followed the protocol recommended by the manufacturer. RT-qPCR was performed with 500 ng total RNA using the Power SYBR-Green Master Mix (cat. no. 4367659) with the 7300 Real-time PCR System (both from Applied Biosystems; Thermo Fisher Scientific, Inc.). Amplification specificity was monitored using the dissociation curve of the amplified product. All data values were normalized with respect to *GUSB*, and the  $\Delta\Delta$ Cq method was used to calculate the fold-change (21). Human Bladder Total RNA (cat. no. AM7990; Applied Biosystems; Thermo Fisher Scientific, Inc.) as control RNA derived from normal bladder tissue.

**Salirasib treatment.** For *in vitro* experiments, salirasib (CAS 162520-00-5; Santa Cruz Biotechnology, Inc., Dallas, TX, USA) was solubilized in 0.1% DMSO. The salirasib/DMSO solution and control vehicle (0.1% DMSO) were prepared in Dulbecco's modified Eagle's medium at different concentrations, and each mixture was placed in cell culture plates (8x10<sup>4</sup>/ml) so that the final salirasib concentrations were 1.6, 3.1, 6.3, 12.5, 25, 50, 100 and 200  $\mu$ M, whereas the DMSO concentration was adjusted to 0.1%. Each cell culture plate was treated with salirasib or control vehicle for 24 h. For *in vivo* experiments, salirasib was solubilized with 0.5% ethanol. The salirasib/ethanol solution was alkalized with 1 N NaOH and then diluted with phosphate buffered saline to yield a 4 mg/ml (pH 8.0) solution. This solution or control vehicle (0.5% ethanol) were intraperitoneally (i.p.) injected daily 100  $\mu$ l per mouse.

**Transfection with small interfering RNA (siRNA).** As described previously (22), T24 and BOY cells were transfected using Lipofectamine RNAiMAX transfection reagent and Opti-MEM (both from Thermo Fisher Scientific, Inc.) together with 10 nM

HRAS siRNA (nos. Hs\_HRAS\_1174, Hs\_HRAS\_1177 and Hs\_HRAS\_1178; Sigma-Aldrich; Merck KGaA) or negative-control siRNA (no. D-001810-10; Thermo Fisher Scientific, Inc.) for loss-of-function experiments. The sequences of the siRNAs were as follows: Hs\_HRAS\_1174\_s, 5'rGUrGrCrCUrGUUrGrGrArCrAUrCrUrGTT; Hs\_HRAS\_1174\_as, 5'rCrArGrGrAUrGUrCrCrArArCrArGrGrCrArCTT; Hs\_HRAS\_1177\_s, 5'rGrArCrGUrGrCrCUrGUUrGrGrArCrAUrCTT; Hs\_HRAS\_1177\_as, 5'rGrAUrGUrCrCrArArCrArGrGrCrArCrGUrCTT; Hs\_HRAS\_1178\_s, 5'rGrGrGrCUUrCrCUrGUrGUrGUUUTT; and Hs\_HRAS\_1178\_as, 5'rArArArCrArCrArCrArGrGrArArGrCrCrCTT. Subsequent experiments were performed 72 h after siRNA transfection.

**Cell proliferation, migration, and invasion assays.** T24 and BOY cells were transfected with 10 nM siRNA by reverse transfection. Cells were seeded in 96-well plates with  $3 \times 10^3$  cells/well for XTT assays. After 72 h, cell proliferation was determined using a Cell Proliferation Kit II (Roche Diagnostics GmbH, Mannheim, Germany) as described previously (22). Cell migration activity was evaluated with wound healing assays. Cells were plated in 6-well plates at  $2 \times 10^5$  cells per well, and after 48 h of transfection the cell monolayer was scraped using a P-20 micropipette tip. The initial (0 h) and residual gap length 18 h after wounding were calculated from photomicrographs as previously described (22). Cell invasion assays were performed using modified Boyden chambers consisting of Matrigel-coated Transwell membrane filter inserts with 8- $\mu$ M pores in 24 well tissue culture plates (BD Biosciences, San Jose, CA, USA). At 72 h after transfection, cells were plated in 24-well plates at  $1 \times 10^5$  cells/well. Minimum essential Eagle's medium containing 10% fetal bovine serum (Equitech-Bio, Inc.) in the lower chamber served as the chemoattractant, as described previously (22). Medium Eagle fetal bovine serum and cells were prepared in the upper chamber and incubated for 24 h.

**Western blot analysis.** Cells were harvested 72 h after transfection, and lysates were prepared in radioimmunoprecipitation assay lysis buffer (Thermo Fisher Scientific, Inc.) containing protease inhibitor cocktail (Sigma-Aldrich; Merck KGaA). Proteins were quantified by Bradford method using BioPhotometer (Eppendorf, Hamburg, Germany). Proteins (50  $\mu$ g) were separated by NuPAGE on 4-12% bis-tris gels (Invitrogen; Thermo Fisher Scientific, Inc.) and transferred to polyvinylidene difluoride membranes. Following blocking in Tris-buffered saline containing 0.1% Tween-20 (TBS-T) with 5% nonfat dry milk for 15 min at 25°C, membranes were washed four times in TBS-T and incubated with primary antibodies overnight at 4°C. Immunoblotting was performed with diluted rabbit polyclonal anti-HRAS antibodies (1:1,000; cat. no. GTX116041; GeneTex, Inc., Irvine, CA, USA), goat polyclonal anti-HIF-1 $\alpha$  antibodies (1:1,000; cat. no. AF1935; R&D Systems, Inc., Minneapolis, MN, USA) and rabbit polyclonal anti- $\beta$ -actin antibodies (1:1,000; cat. no. bs-0061R; BIOSS, Beijing, China) according to the manufacturer's instructions for each antigen. The secondary antibodies were peroxidase-labelled anti-rabbit IgG (1 h at 25°C; 1:5,000; cat. no. 7074S; Cell Signaling Technology, Inc., Danvers, MA, USA) and anti-goat IgG (1 h at 25°C; 1:5,000; cat. no. sc-2020;

Santa Cruz Biotechnology, Inc.). Specific complexes were visualized with an enhanced chemiluminescence detection system (GE Healthcare Life Sciences, Little Chalfont, UK) as described previously (23).

**Proteomic analysis.** To comprehensively investigate metabolic changes in BC cells treated with salirasib, proteomic analysis was performed using iMPAQT (19). Proteins with downregulated expression were detected in salirasib-treated BC cells compared with untreated cells (fold change <0.5) and proteins that were common to both T24 and BOY were identified. The proteins were then categorized into Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways through GeneCodis analysis (genecodis.cnb.csic.es).

**In vivo tumor xenograft model.** To investigate *in vivo* effects of salirasib, a mixture containing 100  $\mu$ l BOY cells ( $5 \times 10^6$ ) and 100  $\mu$ l Matrigel Matrix (Corning Incorporated, Corning, NY, USA) was injected subcutaneously into one side flank of 9 female nude mice (BALB/c nu/nu; 8 weeks old; 16-19 g). The mice were randomly separated into salirasib-treated (n=5) and control (n=4) groups. Each breeding room was kept at a temperature of  $23 \pm 1^\circ\text{C}$  and a humidity of 40-70%. The light/dark cycle was set to 12 h. Food and water was placed to be accessible from each cage. From the day following tumor implantation, salirasib (0.4 mg/mouse, i.p., daily) and control vehicle (0.5% ethanol, 100  $\mu$ l/mouse, i.p., daily) treatment were administered for 25 days. Tumor sizes were measured twice weekly and tumor volumes were calculated as follows: Tumor volume = [(long axis length in millimeters/2) x (short axis length/2)<sup>2</sup> x  $\pi$  x 4]/3. All animal experiments were performed in accordance with institutional guidelines and were approved by the animal care review board of Kagoshima University (Kagoshima, Japan).

**Statistical analysis.** Data are presented as the mean  $\pm$  standard deviation at least three independent experiments. The relationships between two groups were analyzed using Mann-Whitney U tests. The relationships between three or more variables and numerical values were analyzed using Bonferroni-adjusted Mann-Whitney U tests. All analyses were performed using Expert StatView software, version 5.0 (SAS Institute, Inc., Cary, NC, USA). P<0.05 was considered to indicate a statistically significant difference.

## Results

**Expression levels of HRAS in BC and BC cell lines.** The expression levels of HRAS were evaluated using TCGA data from BC samples (n=407) and normal samples (n=19). HRAS expression levels were significantly upregulated in tumor tissues compared with those in normal bladder epithelia (tumor,  $10.314 \pm 0.813$ ; normal,  $9.707 \pm 0.826$ ; P=0.0024, Mann-Whitney U tests; Fig. 1A). Furthermore, HRAS expression was significantly upregulated in patients with BC with mutant HRAS compared with patients with wild-type HRAS (mutant HRAS,  $11.277 \pm 0.805$ ; wild-type HRAS,  $10.267 \pm 0.788$ ; P<0.0001, Mann-Whitney U tests) (Fig. 1B). HRAS mRNA expression was also significantly upregulated in BC cell lines compared to patients with normal bladder tissues (T24,  $5.960 \pm 0.344$ ,

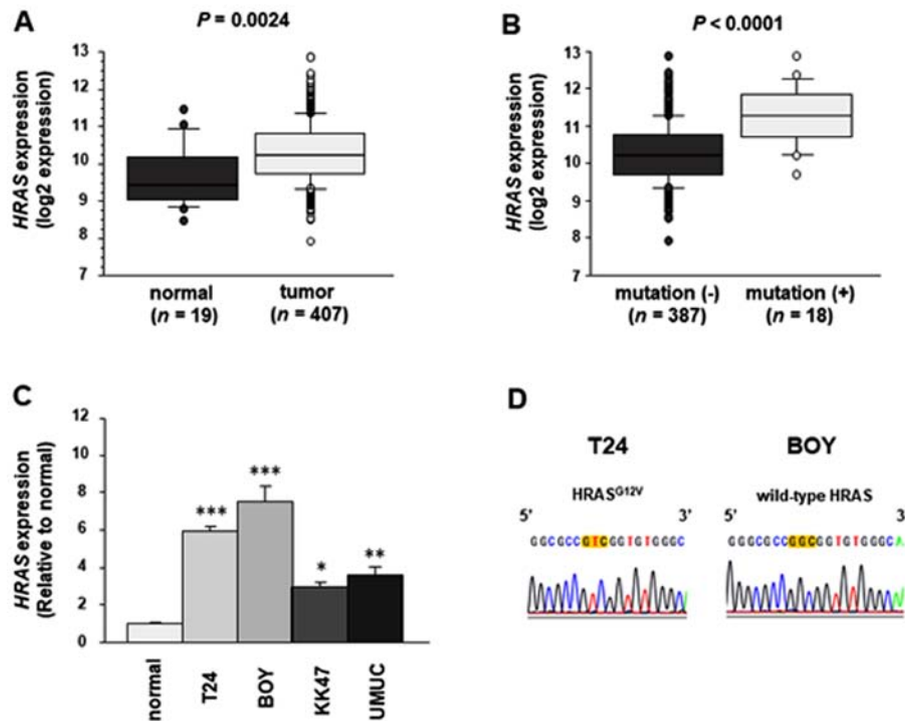


Figure 1. Expression levels of *HRAS* in BC cell lines and clinical BC samples. Significant upregulation of *HRAS* expression levels in reverse transcription-quantitative polymerase chain reaction in (A) BC tumor samples compared with those of normal bladder samples ( $P=0.0024$ ; data analyzed by Mann-Whitney U test are presented as the boxplots) and (B) BC samples with mutant *HRAS* compared with patients with wild-type *HRAS* ( $P<0.0001$ , data analyzed by Mann-Whitney U test are presented as the boxplots) and (C) BC cell lines compared with normal bladder tissues ( $*P=0.0176$ ,  $**P=0.0034$ ,  $***P<0.0001$  vs. normal; data analyzed by Bonferroni-adjusted Mann-Whitney test are presented as the mean  $\pm$  standard deviation from three independent experiments). (D) Difference in *HRAS* mutation status among BC cell lines. Sequencing revealed that T24 had *HRAS*<sup>G12V</sup> (substitution of glycine by valine at codon 12 in *HRAS*) and BOY had wild-type *HRAS*. BC, bladder cancer; *HRAS*, *HRAS* proto-oncogene GTPase.

$P<0.0001$ ; BOY,  $7.528\pm 1.506$ ,  $P<0.0001$ ; KK47,  $2.934\pm 0.464$ ,  $P=0.0176$ ; UMUC,  $3.561\pm 0.854$ ,  $P=0.0034$ ; Bonferroni-adjusted Mann-Whitney U tests; Fig. 1C). Sequencing data from TCGA revealed that T24 cells had *HRAS*<sup>G12V</sup> (the substitution of glycine by valine at codon 12 in *HRAS*) and BOY cells had wild-type *HRAS* (Fig. 1D).

**Effects of *HRAS* knockdown on cell proliferation, migration, and invasion of BC cell lines.** To investigate the functional role of *HRAS* in BC cells, loss-of-function studies were performed using T24 and BOY BC cells transfected with three *si-HRAS* constructs (*si-HRAS-1*, *si-HRAS-2* and *si-HRAS-3*). RT-qPCR analysis and western blot analysis indicated that these siRNAs effectively downregulated *HRAS* mRNA and protein expression in both cell lines (Fig. 2A). XTT assays demonstrated that cell proliferation was inhibited in *si-HRAS* transfectants compared with mock or siRNA-control transfectants (T24, mock  $1.0\pm 0.047$ , control  $1.0\pm 0.015$ , *si-HRAS-1*  $0.701\pm 0.015$ , *si-HRAS-2*  $0.615\pm 0.011$ , *si-HRAS-3*  $0.599\pm 0.024$ ; BOY, mock  $1.0\pm 0.024$ , control  $0.996\pm 0.087$ , *si-HRAS-1*  $0.585\pm 0.030$ , *si-HRAS-2*  $0.499\pm 0.020$ , *si-HRAS-3*  $0.508\pm 0.016$ ; each  $P<0.0001$ , Bonferroni-adjusted Mann-Whitney test; Fig. 2B). Cell migration activity was also significantly inhibited in *si-HRAS* transfectants compared with mock or siRNA-control transfectants (T24, mock  $1.0\pm 0.111$ , control  $1.013\pm 0.117$ , *si-HRAS-1*  $0.684\pm 0.198$ , *si-HRAS-2*  $0.583\pm 0.309$ , *si-HRAS-3*  $0.462\pm 0.348$ ,  $P=0.0019$  and  $P<0.0001$  vs. mock; BOY, mock  $1.0\pm 0.186$ , control  $0.978\pm 0.080$ , *si-HRAS-1*  $0.158\pm 0.105$ , *si-HRAS-2*  $0.438\pm 0.130$ , *si-HRAS-3*  $0.448\pm 0.186$ , each  $P<0.0001$  vs. mock, Bonferroni-adjusted

Mann-Whitney test; Fig. 2C), as was cell invasion activity in Matrigel assays (T24, mock  $1.0\pm 0.280$ , control  $1.443\pm 0.289$ , *si-HRAS-1*  $0.543\pm 0.113$ , *si-HRAS-2*  $0.552\pm 0.154$ , *si-HRAS-3*  $0.353\pm 0.074$ ; BOY, mock  $1.0\pm 0.288$ , control  $1.309\pm 0.268$ , *si-HRAS-1*  $0.117\pm 0.070$ , *si-HRAS-2*  $0.296\pm 0.121$ , *si-HRAS-3*  $0.142\pm 0.086$ , each  $P<0.0001$ , Bonferroni-adjusted Mann-Whitney test; Fig. 2D).

**Effects of salirasib on cell proliferation, migration, and invasion activities in BC cell lines.** The effect of salirasib treatment was investigated in T24 and BOY cells. XTT assays demonstrated that  $\geq 100$   $\mu\text{M}$  salirasib significantly reduced T24 and BOY viability compared with untreated cells (each  $P<0.0001$ , Bonferroni-adjusted Mann-Whitney test; Fig. 3A). Salirasib treatment significantly reduced T24 and BOY cell migration compared with the mock control (T24, mock  $1.0\pm 0.258$ , salirasib  $0.498\pm 0.326$ ,  $P=0.0020$ ; BOY, mock  $1.0\pm 0.470$ , salirasib  $0.556\pm 0.289$ ,  $P=0.0193$ , Mann-Whitney U tests; Fig. 3B) and invasion activity (T24, mock  $1.0\pm 0.336$ , salirasib  $0.200\pm 0.116$ ; BOY, mock  $1.0\pm 0.477$ , salirasib  $0.122\pm 0.125$ ,  $P=0.0009$ , Mann-Whitney U tests; Fig. 3C) relative to untreated cells.

**Proteomic analysis in BC cells treated with salirasib.** To comprehensively investigate metabolic changes in BC cells treated with salirasib, proteomic analysis of metabolism-associated genes was performed using iMPAQT. The results revealed 58 proteins with downregulated expression (fold change  $<0.5$ ) in both T24 and BOY BC cells treated with

Table I. Metabolic changes in bladder cancer cells treated by salirasib.

Gene symbol	Description	Expression ratio (treated/untreated cells)		
		T24	BOY	Mean of T24 and BOY
AHCYL1	Adenosylhomocysteinase-like 1	ND	ND	ND
AK3	Adenylate kinase 3	ND	ND	ND
ALDH3A2	Aldehyde dehydrogenase 3 family, member A2	ND	ND	ND
ALDH9A1	Aldehyde dehydrogenase 9 family, member A1	ND	ND	ND
ASNS	Asparagine synthetase (glutamine-hydrolyzing)	ND	ND	ND
ATP5L	ATP synthase, H <sup>+</sup> transporting, mitochondrial Fo complex, subunit G	ND	ND	ND
ATP6V1E1	Atpase, H <sup>+</sup> transporting, lysosomal, V1 subunit E1	ND	ND	ND
ATP6V1G1	Atpase, H <sup>+</sup> transporting, lysosomal, V1 subunit G1	ND	ND	ND
DCXR	Dicarbonyl/L-xylulose reductase	ND	ND	ND
DERA	Deoxyribose-phosphate aldolase (putative)	ND	ND	ND
DLD	Dihydrolipoamide dehydrogenase	ND	ND	ND
G6PD	Glucose-6-phosphate dehydrogenase	ND	ND	ND
GBA	Glucosidase $\beta$ , acid	ND	ND	ND
GMPPA	GDP-mannose pyrophosphorylase A	ND	ND	ND
GMPR2	Guanosine monophosphate reductase 2	ND	ND	ND
GNPDA1	Glucosamine-6-phosphate deaminase 1	ND	ND	ND
HSD17B12	Hydroxysteroid (17- $\beta$ ) dehydrogenase 12	ND	ND	ND
IDI1	Isopentenyl-diphosphate delta isomerase 1	ND	ND	ND
IVD	Isovaleryl-coa dehydrogenase	ND	ND	ND
MPST	Mercaptopyruvate sulfurtransferase	ND	ND	ND
MTMR1	Myotubularin related protein 1	ND	ND	ND
NDUFS8	NADH dehydrogenase (ubiquinone) Fe-S protein 8, 2 (NADH-coenzyme Q reductase)	ND	ND	ND
NUDT9	Nudix (nucleoside diphosphate linked moiety X)-type motif 9	ND	ND	ND
PAFAH1B3	Platelet-activating factor acetylhydrolase 1b, catalytic subunit 3	ND	ND	ND
PANK4	Pantothenate kinase 4	ND	ND	ND
PFKM	Phosphofructokinase, muscle	ND	ND	ND
PGD	Phosphogluconate dehydrogenase	ND	ND	ND
PGM2	Phosphoglucomutase 2	ND	ND	ND
PLOD3	Procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3	ND	ND	ND
PMVK	Phosphomevalonate kinase	ND	ND	ND
PRIM1	Primase, DNA, polypeptide 1	ND	ND	ND
SCP2	Sterol carrier protein 2	ND	ND	ND
UQCRB	Ubiquinol-cytochrome c reductase binding protein	ND	ND	ND
GAPDH	Glyceraldehyde-3-phosphate dehydrogenase	0.068	0.113	0.091
GOT1	Glutamic-oxaloacetic transaminase 1, soluble (aspartate aminotransferase 1)	ND	0.242	0.121
BCAT1	Branched chain amino-acid transaminase 1, cytosolic	0.247	ND	0.123
AASDHPPT	Amino adipate-semialdehyde dehydrogenase-phosphopantetheinyl transferase	ND	0.256	0.128
UAP1	UDP-N-acetylglucosamine pyrophosphorylase 1	0.107	0.212	0.16
PGM3	Phosphoglucomutase 3	ND	0.393	0.197
PAFAH1B2	Platelet-activating factor acetylhydrolase 1b, catalytic subunit 2	ND	0.405	0.203
BLVRB	Biliverdin reductase B (flavin reductase (NADPH))	ND	0.407	0.204
NDUFS1	NADH dehydrogenase (ubiquinone) Fe-S protein 1, 7 (NADH-coenzyme Q reductase)	ND	0.492	0.246
DLST	Dihydrolipoamide S-succinyltransferase (E2 component of 2-oxo-glutarate complex)	0.293	0.201	0.247
PHGDH	Phosphoglycerate dehydrogenase	0.224	0.284	0.254

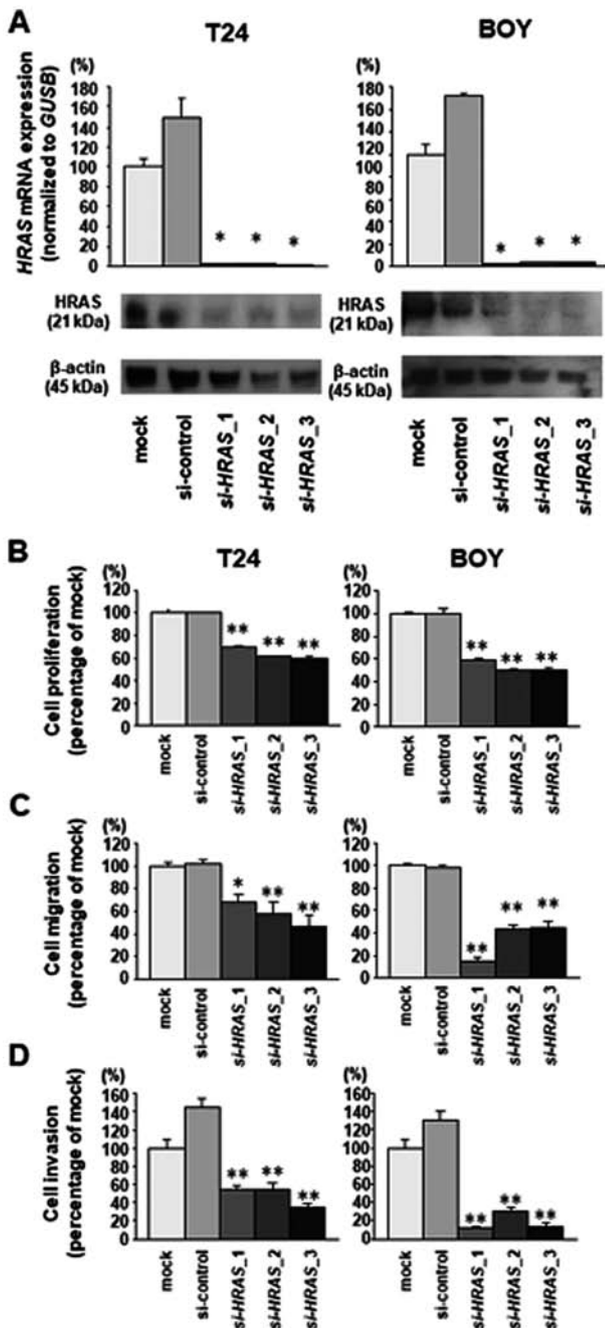


Figure 2. Effects of siRNA-induced *HRAS* knockdown on bladder cancer cell lines. (A) Transfection of T24 and BOY cells with si-*HRAS* show significant ( $P < 0.0001$  vs. mock, Bonferroni-adjusted Mann-Whitney test) downregulation of *HRAS* mRNA and protein expression by reverse transcription-quantitative polymerase chain reaction and western blotting, respectively. Glucuronidase  $\beta$  and  $\beta$ -actin were used as internal and loading controls, respectively. siRNA-induced *HRAS* knockdown inhibited cell proliferation, migration, and invasion as determined by (B) XTT, (C) wound healing and (D) Matrigel invasion assays. ( $P = 0.002$  and  $**P < 0.0001$  vs. mock, Bonferroni-adjusted Mann-Whitney test). All data are presented as the mean  $\pm$  standard deviation. All results represent reproducible data from at least three independent experiments. si, small interfering RNA.

salirasib compared to untreated cells (Table I). GeneCodis analysis to categorize the proteins into KEGG pathways demonstrated that these proteins were included in 50 pathways that were significantly enriched following salirasib treatment (listed in descending order of corrected P-values in Table II; Fig. 4). ‘Oxidative phosphorylation’, ‘pyrimidine

metabolism’, ‘glycolysis/gluconeogenesis’, ‘pentose phosphate pathway’, ‘cysteine and methionine metabolism’, ‘glutathione metabolism’, and ‘purine metabolism’ were significantly downregulated pathways in BC cells treated with salirasib. However, target genes of the RAS effector HIF-1 $\alpha$ , including hexokinase 2, phosphoglycerate kinase 1, pyruvate kinase, muscle (PKM)1, PKM2 and lactate dehydrogenase A, showed only modest downregulation (fold change  $> 0.5$  in T24 and BOY cells) (Table III) (24–26). Furthermore, RT-qPCR analysis and western blot analysis indicated that expression of HIF-1 $\alpha$  was not downregulated in salirasib-treated BC cells (Fig. 5) (5,27).

*Xenograft model study to investigate the in vivo effects of salirasib.* To investigate the *in vivo* effects of salirasib, either salirasib or control vehicle was i.p. injected daily into BC xenograft mice from one day after tumor implantation. There was no difference in tumor growth between the salirasib-treated group ( $n = 5$ ,  $545.9 \pm 187.4$  mm<sup>3</sup>) and control group ( $n = 4$ ,  $511.2 \pm 165.6$  mm<sup>3</sup>) (Fig. 6) on day 27 after tumor implantation.

## Discussion

*HRAS* was the first human oncogene reported in the T24 BC cell line in 1982 (28). Several reports have indicated that *HRAS* mutations critically influence tumorigenesis and development of BC (8,29–33). Haliassos *et al* (29) detected *HRAS* codon 12 point mutations in 66% of BC specimens and the mutant *HRAS* allele in the urine of 47% of patients with BC, Pandith *et al* (33) reported that *HRAS* single nucleotide polymorphism increases BC risk, and rare allele is a predictive marker of advanced bladder tumors. However, RAS had been considered to be ‘undruggable’ because the RAS protein lacked a druggable binding pocket until salirasib was produced. Salirasib inhibits RAS-dependent cell growth by dislodging all isoforms of RAS from the plasma membrane (11,12). The anti-tumor efficacy of salirasib has been demonstrated in several cell lines and xenograft models (17,34–36). Goldberg *et al* (34) demonstrated that salirasib induces pancreatic cancer cell death and tumor shrinkage in mice, and that salirasib was efficient and nontoxic for treatment of glioblastoma in a rat model (35). Charette *et al* (36) reported that salirasib inhibits the growth of hepatocarcinoma cell lines *in vitro* and *in vivo* through RAS and mTOR inhibition. Salirasib was evaluated as a single agent in two clinical trials; however, neither produced promising results in patients with KRAS mutation positive lung adenocarcinoma (13) or refractory hematologic malignancies (14). Even though these clinical trials demonstrated the relative safety of salirasib, diarrhea, nausea and fatigue were the most common toxicities, and there were no grade 4 or 5 drug-associated adverse events or dose-limiting toxicity. On the other hand, results of a combination study of salirasib with gemcitabine to treat pancreatic adenocarcinoma were sufficiently encouraging to warrant further investigation (15,16).

Although the efficacy of salirasib has been reported for several types of cancers, to the best of our knowledge, this is the first report concerning the effect of salirasib in BC. Two BC cell lines were used to evaluate the ability of salirasib to target *HRAS*. T24 carries the *HRAS*<sup>G12V</sup> mutation (substitution of glycine by valine at codon 12 of *HRAS*) and sequencing data demonstrated that BOY cells have wild-type *HRAS*.

Table I. Continued.

Gene symbol	Description	Expression ratio (treated/untreated cells)		
		T24	BOY	Mean of T24 and BOY
HMOX1	Heme oxygenase (decycling) 1	0.165	0.378	0.271
DUT	Deoxyuridine triphosphatase	0.223	0.369	0.296
AHCY	Adenosylhomocysteinase	0.297	0.315	0.306
ATP5A1	ATP synthase, H <sup>+</sup> transporting, mitochondrial F1 complex, $\alpha$ subunit 1, cardiac muscle	0.252	0.363	0.308
SRM	Spermidine synthase	0.297	0.401	0.349
RRM1	Ribonucleotide reductase M1	0.347	0.36	0.353
ACAT1	Acetyl-coa acetyltransferase 1	0.271	0.439	0.355
ISYNA1	Inositol-3-phosphate synthase 1	0.234	0.48	0.357
NDUFA4	NADH dehydrogenase (ubiquinone) 1 $\alpha$ subcomplex, 4	0.286	0.45	0.368
RRM2B	Ribonucleotide reductase M2 B (TP53 inducible)	0.373	0.435	0.404
NUDT2	Nudix (nucleoside diphosphate linked moiety X)-type motif 2	0.443	0.383	0.413
UQCRCQ	Ubiquinol-cytochrome c reductase, complex III subunit VII	0.414	0.451	0.432
CMPK1	Cytidine monophosphate (UMP-CMP) kinase 1, cytosolic	0.462	0.443	0.452
ATP6V1B2	Atpase, H <sup>+</sup> transporting, lysosomal, V1 subunit B2	0.498	0.479	0.489

ND, not detectable (set as 0).

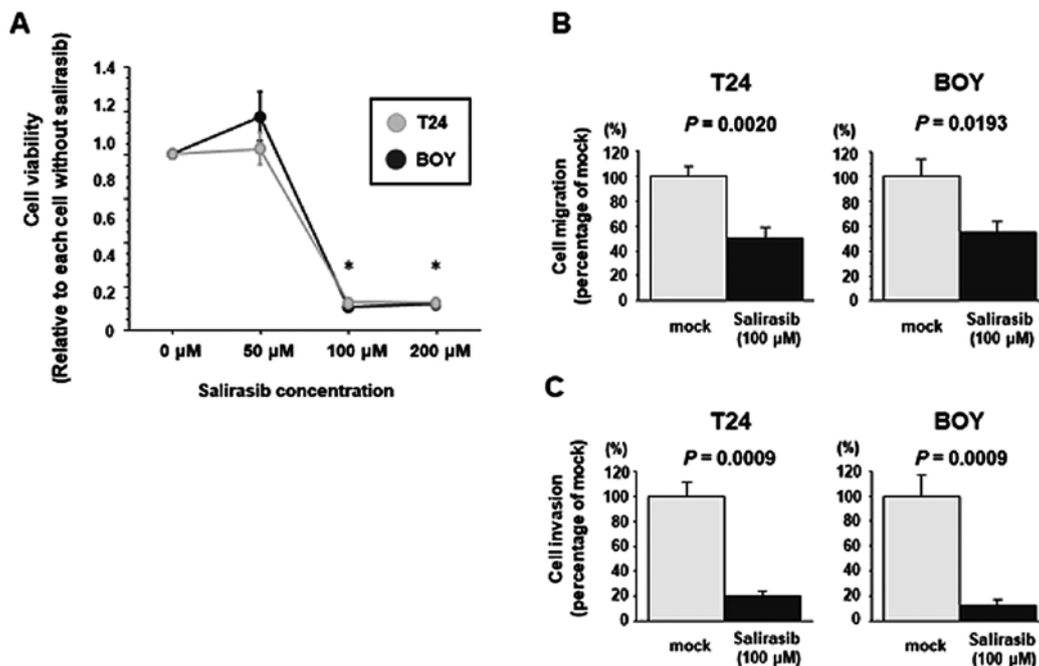


Figure 3. Salirasib inhibition of cell proliferation, migration, and invasion. Compared with vehicle-treated cells, treatment of T24 and BOY cells with 100 and 200  $\mu$ M salirasib inhibited (A) cell proliferation in XTT assays (each  $*P < 0.0001$ , Bonferroni-adjusted Mann-Whitney U test). Treatment of T24 and BOY cells with salirasib (100  $\mu$ M) inhibited (B) cell migration in wound healing assays ( $P = 0.002$  and  $P = 0.0194$ , respectively, Mann-Whitney U test) and (C) cell invasion in Matrigel invasion assays (each  $P = 0.0009$ , Mann-Whitney U test). Data are presented as the mean  $\pm$  standard deviation. All results represent reproducible data from at least three independent experiments.

siRNA-induced *HRAS* knockdown and salirasib inhibition of *HRAS* exerted tumor suppressive effects regardless of *HRAS* mutational status *in vitro*, which was consistent with several previously published results demonstrating a lack of correlation

between *RAS* mutational status and response to *RAS*-targeting therapy (37,38). However, salirasib still required relatively high concentrations to achieve a tumor-suppressive effect *in vitro*, and exhibited no tumor-suppressive effects *in vivo*.

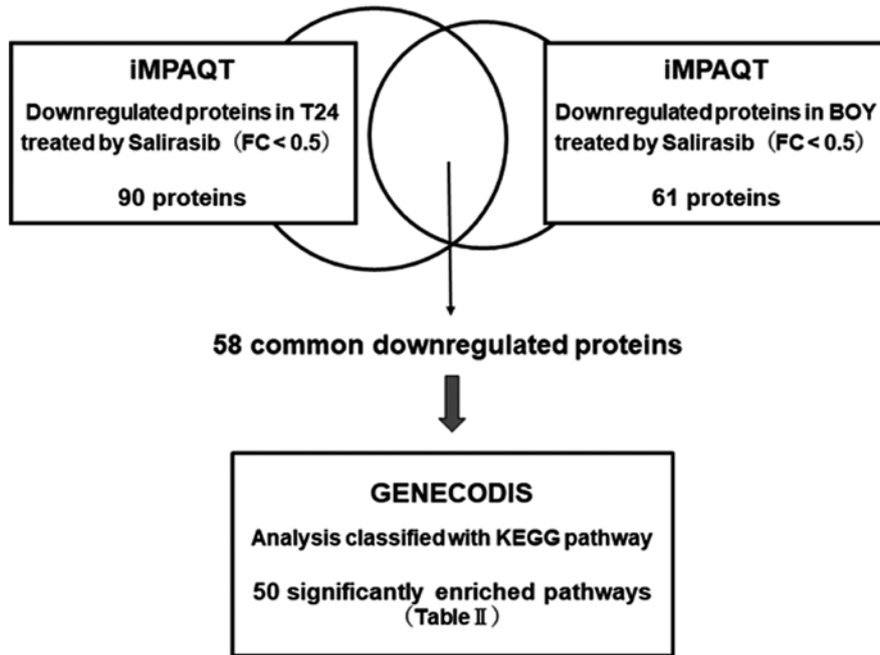


Figure 4. Strategy to identify metabolic pathways downregulated in salirasib-treated BC cells. Proteomic analysis using iMPAQT detected 58 proteins downregulated in both T24 and BOY BC cells treated with salirasib ( $FC < 0.5$ ). Categorization of these proteins into KEGG pathways using GeneCodis analysis showed that 50 pathways were downregulated following salirasib treatment of BC cells. BC, breast cancer; iMPAQT, *in vitro* proteome-assisted multiple reaction monitoring for protein absolute quantification; FC, fold change; KEGG, Kyoto Encyclopedia of Genes and Genomes.

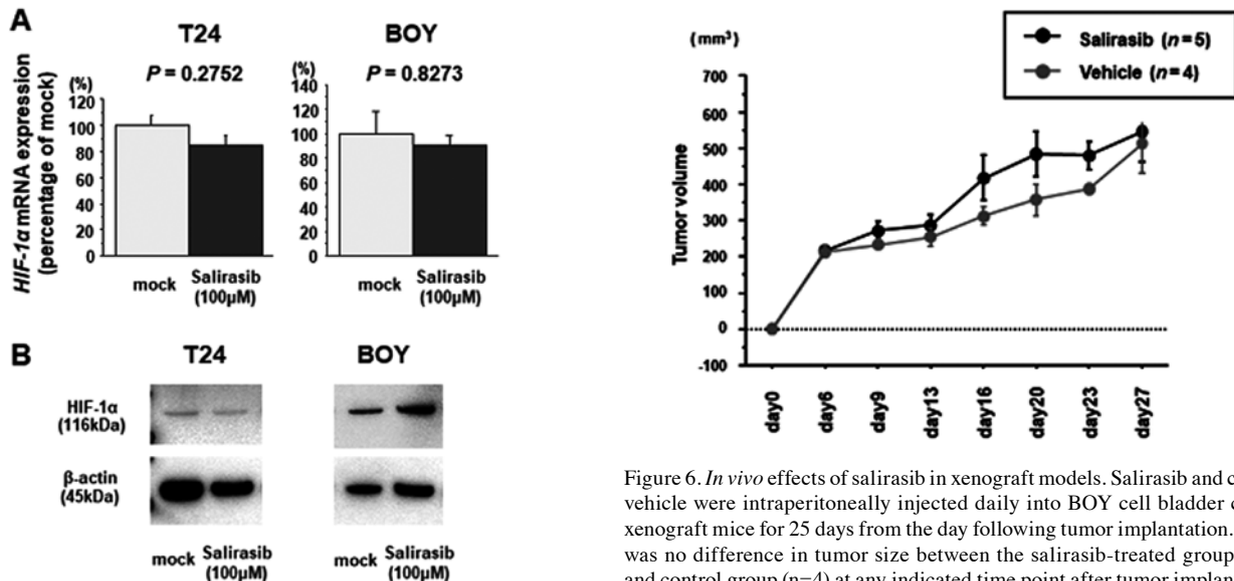


Figure 5. Effect of salirasib on HIF-1 $\alpha$ . (A) Reverse transcription-quantitative polymerase chain reaction analysis and (B) western blot analysis indicated that HIF-1 $\alpha$  mRNA and protein expression were not effectively downregulated in salirasib-treated bladder cancer cells. Data are presented as the mean  $\pm$  standard deviation. All results represent reproducible data from at least three independent experiments. HIF-1 $\alpha$ , hypoxia inducible factor-1 $\alpha$ .

It had been reported that oncogenic RAS predominantly affects the metabolic reprogramming of cancer cells through the upregulation of HIF-1 $\alpha$ , one of target genes of RAS (5). Although salirasib is known to competitively block intracellular signaling via the RAS cascade, there are no reports concerning comprehensive metabolomic analysis of salirasib mechanisms. In the current study, proteomic analysis was performed using

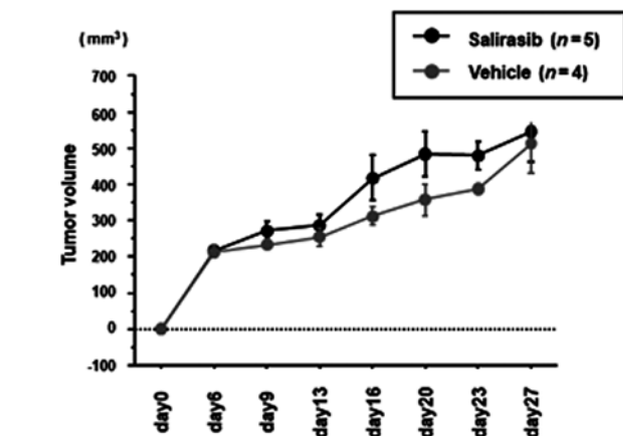


Figure 6. *In vivo* effects of salirasib in xenograft models. Salirasib and control vehicle were intraperitoneally injected daily into BOY cell bladder cancer xenograft mice for 25 days from the day following tumor implantation. There was no difference in tumor size between the salirasib-treated group ( $n=5$ ) and control group ( $n=4$ ) at any indicated time point after tumor implantation. The statistical analysis was performed by Mann-Whitney U test. Data are presented as the mean  $\pm$  standard deviation. The result represents reproducible data from at least three independent experiments.

iMPAQT to investigate metabolic changes in salirasib-treated BC cells. Pathway analysis using the proteomic data indicated that 50 pathways were significantly downregulated following salirasib treatment of BC cells, including 'Oxidative phosphorylation', 'Glycolysis/gluconeogenesis', and 'Pentose phosphate pathway'. However, proteomic analysis showed that the expression of proteins downstream of HIF-1 $\alpha$  were not significantly downregulated. Furthermore, HIF-1 $\alpha$  expression was not efficiently suppressed in salirasib-treated BC cells, although it was previously reported that salirasib suppressed HIF-1 $\alpha$  expression

Table II. Downregulated KEGG pathways in bladder cancer cells treated by salirasib.

KEGG ID	Annotations	Number of genes	Corrected P-value	Genes
00190	Oxidative phosphorylation	10	1.22x10 <sup>-12</sup>	ATP5A1, ATP6V1G1, NDUFS8, ATP5L, ATP6V1E1, NDUFA4, NDUFS1, UQCRB, ATP6V1B2, UQCQRQ
00280	Valine, leucine and isoleucine degradation	6	3.79x10 <sup>-9</sup>	ALDH3A2, DLD, BCAT1, IVD, ALDH9A1, ACAT1
00240	Pyrimidine metabolism	7	4.84x10 <sup>-9</sup>	CMPK1, NUDT2, RRM2B, AK3, RRM1, PRIM1, DUT
00010	Glycolysis/gluconeogenesis	6	1.29x10 <sup>-8</sup>	ALDH3A2, PGM2, DLD, ALDH9A1, GAPDH, PFKM
00030	Pentose phosphate pathway	5	1.43x10 <sup>-8</sup>	PGM2, DERA, G6PD, PGD, PFKM
00270	Cysteine and methionine metabolism	5	4.39x10 <sup>-8</sup>	MPST, AHCY, AHCYL1, SRM, GOT1
00480	Glutathione metabolism	5	8.35x10 <sup>-8</sup>	RRM2B, G6PD, RRM1, SRM, PGD
00230	Purine metabolism	7	8.57x10 <sup>-8</sup>	NUDT2, PGM2, GMPR2, RRM2B, RRM1, NUDT9, PRIM1
05010	Alzheimer's disease	7	9.31x10 <sup>-8</sup>	ATP5A1, NDUFS8, NDUFA4, NDUFS1, UQCRB, GAPDH, UQCQRQ
00520	Amino sugar and nucleotide sugar metabolism	5	1.18x10 <sup>-7</sup>	PGM2, UAP1, GMPPA, PGM3, GNPDA1
00310	Lysine degradation	5	1.19x10 <sup>-7</sup>	ALDH3A2, ALDH9A1, PLOD3, DLST, ACAT1
05012	Parkinson's disease	6	4.55x10 <sup>-7</sup>	ATP5A1, NDUFS8, NDUFA4, NDUFS1, UQCRB, UQCQRQ
05016	Huntington's disease	6	2.79x10 <sup>-6</sup>	ATP5A1, NDUFS8, NDUFA4, NDUFS1, UQCRB, UQCQRQ
00620	Pyruvate metabolism	4	2.79x10 <sup>-6</sup>	ALDH3A2, DLD, ALDH9A1, ACAT1
00330	Arginine and proline metabolism	4	8.56x10 <sup>-6</sup>	ALDH3A2, ALDH9A1, SRM, GOT1
00900	Terpenoid backbone biosynthesis	3	8.57x10 <sup>-6</sup>	PMVK, IDI1, ACAT1
00770	Pantothenate and CoA biosynthesis	3	1.13x10 <sup>-5</sup>	BCAT1, AASDHPPT, PANK4
00410	β-alanine metabolism	3	3.82x10 <sup>-5</sup>	ALDH3A2, ALDH9A1, SRM
04966	Collecting duct acid secretion	3	3.82x10 <sup>-5</sup>	ATP6V1G1, ATP6V1E1, ATP6V1B2
00640	Propanoate metabolism	3	6.87x10 <sup>-5</sup>	ALDH3A2, ALDH9A1, ACAT1
00051	Fructose and mannose metabolism	3	9.37x10 <sup>-5</sup>	GMPPA, MTMR1, PFKM
00071	Fatty acid metabolism	3	0.000127	ALDH3A2, ALDH9A1, ACAT1
00380	Tryptophan metabolism	3	0.000127	ALDH3A2, ALDH9A1, ACAT1
05110	Vibrio cholerae infection	3	0.000264	ATP6V1G1, ATP6V1E1, ATP6V1B2
05120	Epithelial cell signaling in <i>Helicobacter pylori</i> infection	3	0.000509	ATP6V1G1, ATP6V1E1, ATP6V1B2
05323	Rheumatoid arthritis	3	0.000953	ATP6V1G1, ATP6V1E1, ATP6V1B2
00053	Ascorbate and aldarate metabolism	2	0.002054	ALDH3A2, ALDH9A1
00052	Galactose metabolism	2	0.002137	PGM2, PFKM
00340	Histidine metabolism	2	0.002381	ALDH3A2, ALDH9A1
00020	Citrate cycle (TCA cycle)	2	0.002464	DLD, DLST
00040	Pentose and glucuronate interconversions	2	0.002546	ALDH3A2, DCXR
00250	Alanine, aspartate and glutamate metabolism	2	0.002548	ASNS, GOT1
00260	Glycine, serine and threonine metabolism	2	0.002548	DLD, PHGDH
00565	Ether lipid metabolism	2	0.002791	PAFAH1B3, PAFAH1B2
04145	Phagosome	3	0.002924	ATP6V1G1, ATP6V1E1, ATP6V1B2
00860	Porphyrin and chlorophyll metabolism	2	0.004011	HMOX1, BLVRB
00561	Glycerolipid metabolism	2	0.004873	ALDH3A2, ALDH9A1
04260	Cardiac muscle contraction	2	0.011811	UQCRB, UQCQRQ

Table II. Continued.

KEGG ID	Annotations	Number of genes	Corrected P-value	Genes
04146	Peroxisome	2	0.011811	PMVK, SCP2
00400	Phenylalanine, tyrosine and tryptophan biosynthesis	1	0.013097	GOT1
00072	Synthesis and degradation of ketone bodies	1	0.022922	ACAT1
00290	Valine, leucine and isoleucine biosynthesis	1	0.024264	BCAT1
04122	Sulfur relay system	1	0.024264	MPST
00740	Riboflavin metabolism	1	0.026063	BLVRB
00360	Phenylalanine metabolism	1	0.036111	GOT1
00120	Primary bile acid biosynthesis	1	0.036111	SCP2
00511	Other glycan degradation	1	0.03752	GBA
00630	Glyoxylate and dicarboxylate metabolism	1	0.038867	ACAT1
01040	Biosynthesis of unsaturated fatty acids	1	0.044309	HSD17B12
00910	Nitrogen metabolism	1	0.04748	ASNS

KEGG, Kyoto Encyclopedia of Genes and Genomes.

Table III. Effects of salirasib downstream of hypoxia inducible factor-1 $\alpha$ .

Name	Description	T24 fold change	BOY fold change	Mean of T24 and BOY
PKM2	Pyruvate kinase, muscle 2	1.369	1.194	1.281
LDHA	Lactate dehydrogenase A	0.918	1.005	0.961
PKM1	Pyruvate kinase, muscle 1	0.900	1.044	0.972
PGK1	Phosphoglycerate kinase 1	0.857	1.192	1.025
HK2	Hexokinase 2	0.735	0.697	0.716
ENO1	Enolase 1, ( $\alpha$ )	0.453	0.587	0.520
GAPDH	Glyceraldehyde-3-phosphate dehydrogenase	0.068	0.113	0.091

in other type of cancer cells (27). Therefore, downregulation of RAS target genes in *in vitro* assays involving BC cell lines may require a high concentration of salirasib, and this need for high concentrations was responsible for the lack of tumor suppressive effects observed in the BC xenograft mouse model. In this study, whether factors downstream of HIF-1 $\alpha$  were insufficiently downregulated in the tumors from animal experiments was not analyzed by iMPAQT, because iMPAQT is so sensitive that contamination of surrounding tissues adjacent to tumor tissue may make the interpretation of the results difficult. However, these analyses of micro-dissected *in vivo* samples will be performed in the future. Recently, a novel RAS inhibitor developed using an innovative approach was reported to inhibit tumor growth in animal models of RAS-dependent cancers at low concentrations (39). This novel RAS inhibitor was computationally designed to target multiple sites on RAS proteins, thus enabling sufficient affinity and selectivity for pharmacological RAS inhibition. This new inhibitor may provide successful targeting of RAS in the near future. Therefore, clinical trials with these inhibitors or next-generation RAS inhibitors are required to improve cancer treatment options in the near future.

In conclusion, the current study demonstrated that salirasib and siRNA-induced *HRAS* knockdown produced tumor suppressive effects regardless of *HRAS* mutational status in BC cell lines. However, high concentrations of salirasib were required to inhibit cell proliferation, migration and invasion activity *in vitro*, and the same high concentrations exhibited no tumor suppressive effects *in vivo*. Proteomic analysis revealed that several metabolic pathways were significantly downregulated in BC cells treated with salirasib. However, salirasib treatment of BC cells did not significantly affect expression of genes targeted by HIF-1 $\alpha$  in BC cells. These findings provide novel information concerning the mechanism of salirasib effects, and suggest that novel therapeutics involving combination therapies of salirasib with other inhibitors, or the newly-identified novel RAS inhibitor, may be effective for treating BC and other types of cancer.

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### Availability of data and materials

The analyzed datasets generated during the study are available from the corresponding author on reasonable request.

### Authors' contributions

HY conceived and designed the experiments. SS, HY, KM, MY, TS and YO performed the experiments. SS, HE, HY and KM performed the validation and formal analysis. SS and KM wrote the manuscript. HE, HY and MN interpreted experimental data for the work, and reviewed and revised the manuscript. HE, HY and MN acquired funding. All authors have read and approved the final manuscript.

### Ethics approval and consent to participate

Not applicable.

### Patient consent for publication

Not applicable.

### Competing interests

The authors declare that they have no competing interests.

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