tumor activity remain elusive. To date, factors that have been identified as correlated with the efficacy of sorafenib include phosphorylated extracellular signal-regulated kinase 1 (p-ERK) (6), serum des- γ -carboxyprothrombin level (7), phosphorylated c-Jun protein (8), and fibroblast growth factor-3/4 gene amplification (3), but their clinical utility as predictive biomarkers has not been established.

In the present study, we developed a new technique, high-density fluorescence reverse-phase protein array (RPPA), and used it to search for a biomarker that would identify patients in whom sorafenib would be effective, employing a large library of phosphorylation-site-specific antibodies. RPPA represents an emerging technology for proteomics, and it is well suited for the profiling of phosphorylated proteins. It involves micro-format dot immunoblotting of lysates from tissues or cells (9), allowing simultaneous monitoring of the expression of a particular phosphoprotein in hundreds to thousands of samples under identical conditions in a highly quantitative manner (10). In this study we profiled the activation status of 180 key signaling nodes across a panel of 23 HCC cell lines and identified *de novo* activation of mTOR signaling in sorafenib-resistant HCC cells.

EXPERIMENTAL PROCEDURES

Cell Lines and Antibodies—Cell lines used for generating the cancer cell line RPPA are listed in supplemental Table S1 and were maintained according to their suppliers' recommendations. Recombinant EGF was obtained from R&D Systems (Minneapolis, MN). A total of 180 phosphorylation-site-specific antibodies and their dilutions used for RPPA analysis are listed in supplemental Table S2. The specificity of each antibody was verified by immunoblotting or had been previously described by other investigators.

RPPA—Cells were collected by scraping and stored at —801°C until use. Cell lysates were prepared with RIPA buffer (Thermo Scientific, Rockford, IL) supplemented with phosphatase (Thermo Scientific) and protease (Sigma, St. Louis, MO) inhibitor cocktails. Protein concentrations of lysates were determined via the Bradford method (Bio-Rad Laboratories, Hercules, CA). The lysates were serially diluted 2-fold four times and printed in quadruplicate onto ProteoChip glass slides (Proteogen, Seoul, South Korea) using a robotic spotter (Genex Arrayer, Kaken Genegs Inc., Chiba, Japan).

The RPPA slides were incubated overnight with primary antibodies. Following tyramide signal amplification (Dako Cytomation, Glostrup, Denmark), streptavidin Alexa Fluor 647 conjugate (Invitrogen, Carlsbad, CA) was applied to the slides (11). Fluorescence images were captured by an InnoScan 700 microarray scanner (Innopsys, Carbonne, France) and quantified using Mapix software (Innopsys). After background subtraction, values relative to γ -tubulin were subjected to quantile normalization (12) to ensure a uniform distribution of values for each slide in a set of slides. Unsupervised hierarchical clustering, using the Euclidean metric and Ward's method, was conducted with R 2.13.0. The signaling components of the mTOR and MAPK pathways were selected based on KEGG pathway maps and used for clustering analyses.

Immunoblot Analysis — Immunoblot analyses were performed using the NuPAGE Bis-Tris or Tris-Acetate electrophoresis system (Invitrogen) as described previously (13). All antibodies except for an antip-RSK (S380) antibody (R&D Systems) were obtained from Cell Signaling Technology (Danvers, MA). Signals were detected with the

ImageQuant LAS 4010 system (GE Healthcare, Giles, UK) and quantified using the ImageQuant TL software package (GE Healthcare).

Growth Inhibition Assay—Sorafenib, RAD001 (everolimus), and SL0101 were purchased from Toronto Research Chemicals Inc. (North York, Ontario, Canada). Cl-1040 and AZD8055 were from Selleck Chemicals (Houston, TX). Stock solutions of the chemicals were prepared in dimethyl sulfoxide and stored at $-20\mbox{l}^{\circ}\text{C}$ until use. Cells were seeded into 96-well cell culture plates in triplicate at a density of 3000 cells per well. On the following day, serially diluted drugs were added, and 72 h later cell viability was measured using the CellTiter-Glo Luminescent Cell Viability Assay (Promega, Fitchburg, WI). Relative cell viability was calculated as a percentage of a control treated with 0.1% dimethyl sulfoxide after background subtraction. All experiments were repeated at least three times. The data were modeled using a four-parameter log-logistic nonlinear regression curve fit with a sigmoid dose response. These curves were drawn using R 2.13.0, and IC_{50} values were calculated accordingly.

Immunohistochemistry—Formalin-fixed, paraffin-embedded sections of needle biopsy samples obtained from nine HCC patients before administration of sorafenib at Wan Fang Hospital and the Taipei Medical University Hospital were immunostained with anti-p-RPS6 Ser235/236 (#2211, Cell Signaling Technology) or anti-RPS6 (#2217, Cell Signaling Technology) antibody, as described previously (13). The stained slides were evaluated by pathologists and classified according to the percentage of positively stained cells (0 = 0%, 1 = 1% to 25%, 2 = 26% to 50%, 3 = 51% to 75%, and 4 = 76% to 100%) and the intensity of staining (0, absent; 1, weak; 2, moderate; and 3, strong). A specimen was defined as positive when either the percentage of positively stained cells or the intensity of staining was 3 or higher. The use of clinical materials was approved by the respective institutional review boards.

Kinome Sequencing-Genomic DNA was extracted from 20 HCC cell lines using the DNeasy Blood and Tissue kit (Qiagen, Hilden, Germany), in accordance with the manufacturer's protocol. DNA concentration was determined using a NanoDrop 2000 spectrophotometer (Thermo Scientific). Three micrograms of genomic DNA was used to construct libraries for sequencing. The quality of the constructed libraries was assessed using an Agilent 2100 Bioanalyzer (Agilent Technologies, Santa Clara, CA). All the exon and 5'- and 3'-flanking sequences (200 bp) of 511 kinase genes were captured using a customized SureSelect Target Enrichment System (Agilent Technologies) according to the Illumina Paired-End Sequencing Platform Library Prep Protocol Version 1.0 (Agilent Technologies). Captured DNA fragments (~300 bp) were sequenced using a Genome Analyzer IIx sequencer (Illumina, San Diego, CA). Base calling was performed using the Illumina Pipeline (v1.4) with default parameters. Only paired end (2 \times 75 bases) sequence reads that passed the quality control were mapped to the human reference genome build hg19 (UCSC hg19) using BWA (14) with default parameters. Sequencing artifacts were eliminated using Picard MarkDuplicates. Variants were called with SAMtools (15) and annotated using Annovar ENREF 25 (16). The final set of novel variant calls was identified using the following thresholds: SNP quality ≥ 228, coverage ≥ 20 reads, frequency ≥ 10%, and not deposited in the dbSNP database (www.ncbi.nlm. nih.gov/projects/SNP/) (version 135).

Evaluation of Synergistic Drug Combinations—The synergistic interaction of drug combinations was evaluated using the Chou–Talalay median-dose effect method (17) with CompuSyn software. AZD8055 and Cl-1040 were mixed at the ratio of their IC_{50} values (1:300). The mixed solution was 2-fold serially diluted five times and added to the cells. Combination Index values were calculated at the points causing 50%, 75%, and 90% reduction of cell viability. Combination Index values equal to 1, >1, and <1 indicate additive, antagonistic, and synergistic interactions, respectively.

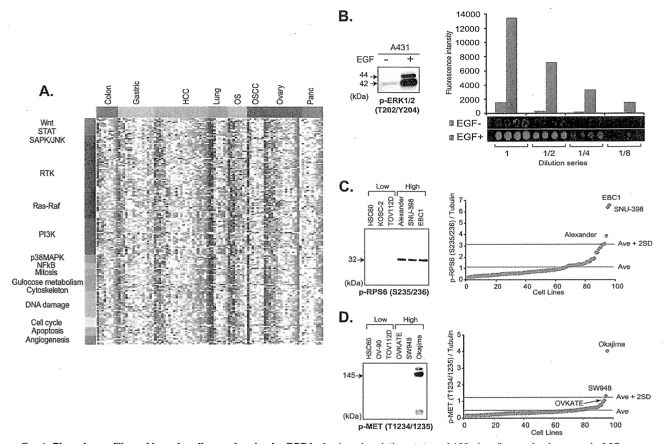


Fig. 1. **Phosphoprofiling of key signaling molecules by RPPA.** *A*, phosphorylation status of 180 signaling nodes in a panel of 95 cancer cell lines cultured in the presence of 10% FCS. Red and blue colors indicate high- and low-level phosphorylation, respectively. STAT, signal transducers and activators of transcription; SAPK/JNK, stress-activated protein kinase/c-Jun NH₂-terminal kinase; RTK, receptor tyrosine kinase; Pl3K, phosphatidylinositol 3′-kinase; MAPK, mitogen-activated protein kinase; NFκB, nuclear factor-kappaB; OS, osteosarcoma; OSCC, oral squamous cell carcinoma. *B*, immunoblot (*left*) and RPPA (*right*) analyses of A431 cells cultured without (–) and with (+) EGF for 10 min with anti-p-ERK1/2 (T202/Y204) antibody. The mean fluorescence intensity in arbitrary units (*top*) and images (*bottom*) of quadruplicate RPPA spots of lysate undiluted (1) and diluted 1:2 (1/2), 1:4 (1/4), and 1:8 (1/8) -fold are shown (*right*). *C*, *D*, relative p-RPS6 S235/236 (*C*) and p-Met T1234/1235 (*D*) expression of 95 cell lines determined via RPPA (*right*). Cell lines with the three highest and three lowest levels of expression were selected and subjected to immunoblotting with the same antibody (*left*). Ave, average.

RESULTS

Generation of the High-density RPPA and Phosphoprotein Profiling—We constructed an RPPA onto which lysates of 95 cell lines derived from eight different types of cancer (listed in supplemental Table S1) cultured in the presence and absence of 10% fetal calf serum (FCS) for 17 h and A431 cells untreated or treated with 200 ng/ml EGF for 10 min were randomly plotted. Each lysate was serially diluted (1:1, 1:2, 1:4, and 1:8) and spotted in quadruplicate (16 spots per lysate). This level of high-density spotting (3072 samples per array slide) was achievable because of the highly hydrophobic surface of the array slides, which prevented diffusion of the protein samples.

By applying 180 phosphorylation-site-specific antibodies (listed in supplemental Table S2), we determined the activation status of signaling proteins (Fig. 1A). A lysate of A431 cells treated with EGF was included as a positive internal

control. A431 cells carry amplification of the *EGFR* (EGF receptor) gene. We confirmed that a >6-fold increase in the signal intensity of ERK1/2 proteins phosphorylated at the threonine 202/tyrosine 204 residue (p-ERK1/2 T202/Y204) was detectable after treatment with EGF (Fig. 1*B*).

To further verify the data obtained via RPPA, lysates of representative cell lines were electrophoresed and blotted with the same antibodies. In the RPPA analysis, EBC1, SNU-398, and Alexander cells showed a high signal intensity for anti-p-RPS6 S235/236 antibody, exceeding the average plus 2 S.D. for 96 cell lines, whereas TOV112D, KOSC-2, and HSC60 cells showed a signal intensity below the average (Fig. 1C, right). The results we obtained from immunoblotting were consistent (Fig. 1C, left).

The glass slides that we used for construction of the RPPA were free of any autofluorescence noise. The use of fluorescent dyes and original signal enhancement significantly in-

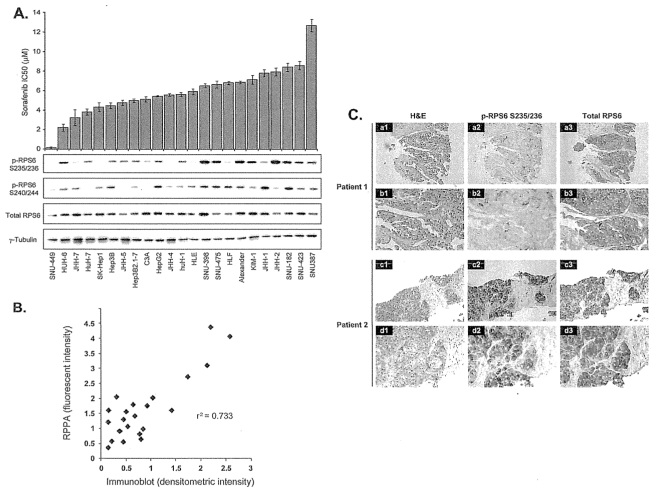


Fig. 2. **p-RPS6 S235/236 correlates with the sensitivity of HCC to sorafenib.** *A, upper graph,* IC_{50} values for sorafenib against 23 HCC cell lines sorted from the most sensitive (*left*) to resistant (*right*) ones. Columns and error bars represent the mean and S.D. of three independent experiments, respectively. *Lower panels,* immunoblot analysis of pRPS6 S235/236, pRPS6 S240/244, RPS6, and γ -tubulin (loading control) expression in the 23 HCC cell lines. *B,* correlation between RPPA and immunoblot analyses of p-RPS6 S235/236 expression in the 23 HCC cell lines ($R^2 = 0.733$). *C,* detection of p-RPS6 S235/236 in pretreatment biopsy samples. Hematoxylin and eosin (H&E) (a-d1) and immunoperoxidase staining with anti-p-RPS6 S235/236 (a-d2) and total RPS6 (a-d3) antibodies of HCC biopsy specimens obtained from a responder (patient 1 (a and b)) and a representative non-responder (patient 2 (c and d)) to sorafenib. Original magnification was ×40 (a1–3 and c1–3) and ×200 (b1–3 and d1–3).

creased the sensitivity of signal detection. In fact, MET protein with a high level of phosphorylation (p-MET T1234/1235) in Okajima cells was detectable via immunoblotting, whereas the MET protein with a relatively low level of phosphorylation in SW948 and OVKATE cells was undetectable (Fig. 1D).

p-RPS6 S235/236 Correlates with the Sensitivity of HCC Cells to Sorafenib—The cancer cell protein array contained 23 HCC cell lines exhibiting a wide variety of sensitivities to sorafenib (Fig. 2A, upper portion). SNU-449 was the most sensitive, with a half-maximal (50%),inhibitory concentration (IC₅₀) of 0.172 μ M. It was ~70-fold more sensitive than the least sensitive cell line, SNU-387 (IC₅₀ = 12.68 μ M). We then compared the IC₅₀ value of each HCC cell line with the phosphorylation level of 180 signaling nodes. Spearman's correla-

tion coefficient analysis (supplemental Table S3) revealed that p-RPS6 S235/236 had the highest positive correlation (r=0.58, p=0.0044), followed by p-RPS6 at the serine 240/244 residues (p-RPS6 S240/244) (r=0.55, p=0.0070). 90-kDa ribosomal S6 kinase 2 (RSK2) protein phosphorylated at the serine 227 residue showed the third most significant correlation. RSK2 is one of the enzymes that phosphorylate RPS6 (18).

Consistent with the RPPA data, intense signals for p-RPS6 S235/236 were detected in the sorafenib-resistant cell lines via immunoblotting (Fig. 2A, lower portion). The quantified immunoblot data correlated well with those of RPPA ($r^2 = 0.733$), thus confirming the precision of the RPPA (Fig. 2B). p-RPS6 S235/236 and p-RPS6 S240/244 exhibited different

phosphorylation patterns among several cell lines (e.g. HLF, KIM1, JHH-1, and JHH-2) (Fig. 2A, lower portion), suggesting that phosphorylation of S235/236 and S240/244 residues may be mediated by distinct regulatory processes.

p-RPS6 S235/236 Is a Potential Predictor of Response to Sorafenib-We next evaluated whether high levels of p-RPS6 S235/236 were indicative of HCC resistance to sorafenib in clinical samples (supplemental Table S4). Expression of p-RPS6 S235/236 was examined in biopsy specimens collected from nine HCC patients prior to sorafenib treatment (400 mg twice a day). Eight patients showed intense staining for p-RPS6 (Fig. 2C). Four patients (Cases 4, 6, 7, and 9) with p-RPS6-positive tumors discontinued sorafenib treatment because of disease progression within 2.3 months. Four patients (Cases 2, 3, 5, and 8) died as a result of disease progression after starting sorafenib treatment. In contrast, the remaining patient (Case 1), whose tumor was negative for p-RPS6, received sorafenib for 24 months and survived for 27 months. In this particular patient, tumor regression was confirmed by computed tomography scans performed three months after sorafenib administration and remained stable for another three months. In addition, the α -fetoprotein level dropped from 1621 to 314 ng/ml and remained low for 10 months. These results provide preliminary evidence that that high expression of p-RPS6 S235/236 might be useful for predicting which patients are unlikely to respond to sorafenib. As biopsy is not performed routinely before sorafenib treatment, we were unable to further validate the clinical significance of p-RPS6 S235/236 by examining additional cases.

mTOR Pathway Activation in Sorafenib-resistant Cells—Given the association between p-RPS6 and sensitivity of HCC cell lines to sorafenib, we assessed the effects of sorafenib on p-RPS6 in representative sorafenib-sensitive and -resistant cell lines. The sorafenib-sensitive HUH-6 and HuH-7 cell lines demonstrated substantial dose-dependent decreases in p-RPS6 levels following treatment with sorafenib (Fig. 3A). Sorafenib also diminished the phosphorylation of downstream molecules in the MAPK pathway, ERK and RSK, in a dose-dependent manner (Fig. 3A), implying that the reduction of p-RPS6 in sorafenib-sensitive cells was likely attributable to blockade of the MAPK pathway (Fig. 3B).

In contrast, the phosphorylation of RPS6 S235/236 in sorafenib-resistant cell lines, especially JHH-2, SNU-423, and SNU-387 cells, was insensitive to the same sorafenib treatment (Fig. 3C). The level of p-RPS6 in JHH-1 and SNU-182 cells decreased to some extent after sorafenib treatment, but a high concentration (10 μ M) of sorafenib was necessary in order to suppress the phosphorylation of RPS6 completely (Fig. 3C), reflecting that the regulation of p-RPS6 in sorafenib-resistant cell lines is different from that in sensitive cell lines. RPS6 is also known to be phosphorylated by 70-kDa ribosomal S6 kinases (S6K) downstream of mTOR (Fig. 3B) (19, 20). We therefore speculated that the mTOR pathway might be alternatively activated in sorafenib-resistant cell lines. In

fact, we found that sorafenib-resistant cells had a high level of p-S6K1 (Fig. 3C), whereas p-S6K1 was barely detectable in sorafenib-sensitive cells (Fig. 3A).

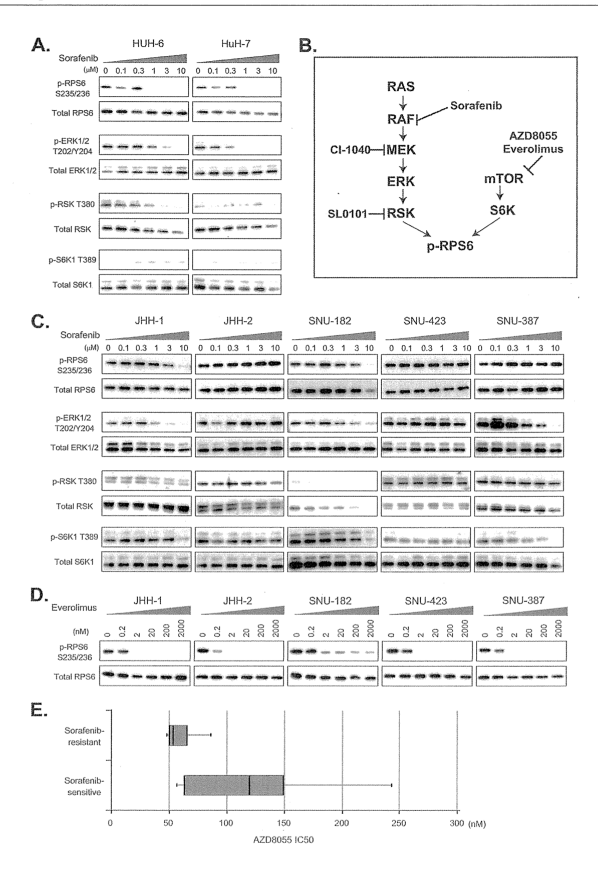
The phosphorylation of ERK in JHH-1 and SNU-182 was suppressed to some extent by sorafenib, but the low level of p-RSK and high level of p-S6K1 indicate that the main regulator of RPS6 phosphorylation was mTOR signaling rather than MAPK signaling.

Absence of Genetic Alterations in the MAPK and mTOR Pathways—RAF kinases are among the known targets of sorafenib, but sorafenib-resistant JHH2, SNU-423, and SNU-387 cells exhibited sustained activation of molecules located downstream of RAF (ERK and RSK), even in the presence of sorafenib (Fig. 3C), suggesting sorafenib-insensitive activation of the MAPK pathway.

To clarify the molecular mechanism driving the activation of the mTOR and MAPK pathways in sorafenib-resistant cells, we sequenced the entire exons of 511 kinases (listed in supplemental Table S5) in 20 HCC cell lines using a next-generation sequencer. Supplemental Table S6 lists all of the 322 genetic alterations that were not deposited in the dbSNP database. Due to the unavailability of normal counterparts, we were unable to determine whether these alterations were somatic. Eight kinds of DNA alterations were evident in the known mTOR and MAPK pathway genes (Supplemental Table S7). b-RAF V600E, found in SK-Hep1 cells, is a known driver mutation frequently observed in malignant melanoma (21). Two kinds of alterations were identified in the ATP-binding (S72A (JHH-7)) and AGC-kinase C-terminal (K335T (huH-1, SNU-475, and SNU-185)) domains of the RSK1 genes. Three kinds of alterations were found in the proline-rich domain (A420V (11 cell lines including SNU449) and V422I (HUH-6)) and catalytic (P267L (SK-Hep1, JHH-4, Kim1, and JHH-1)) domains of the S6K2 gene.

These eight alterations were validated using a conventional sequencing method (Supplemental Table S7), but no genetic alteration was specific to sorafenib-resistant cell lines. Infrequent alteration of the mTOR and MAPK pathway genes in HCC has been demonstrated by conventional sequencing analysis of surgical samples (22), and this was consistent with the present comprehensive sequencing data. The aberrant activation of the mTOR and MAPK pathways in sorafenib-resistant HCC cells was likely attributable to complex interplay between other oncogenic and anti-oncogenic pathways, or post-translational modifications.

mTOR Inhibitors Repress the Proliferation of Sorafenib-resistant Cells—The marked inhibition of p-RPS6 S235/236 following exposure to an mTOR inhibitor, everolimus, at a concentration as low as 2 nm (Fig. 3D) confirmed that activation of the mTOR pathway is responsible for the sorafenib-insensitive phosphorylation of RPS6 S235/236 in sorafenib-resistant cells. Consistently, sorafenib-insensitive cells tended to be more sensitive to another mTOR inhibitor,



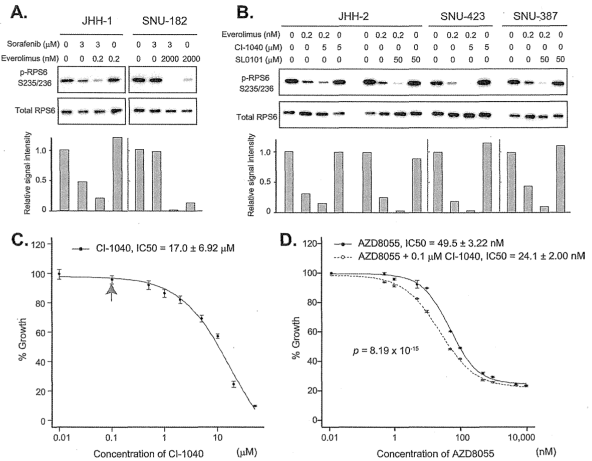


Fig. 4. **Synergy of mTOR and MAPK inhibitors.** A, sorafenib-resistant JHH-1 and SNU-182 cells were treated with the indicated concentrations of sorafenib and everolimus, and the expression of p-RPS6 S235/236 and total RPS6 was examined via immunoblotting (top). The bottom panel indicates intensity relative to control blots (no treatment). B, sorafenib-resistant JHH2, SNU-423 and SNU-387 cells were treated with the indicated concentrations of Cl-1040, SL0101, and everolimus, and the expression of p-RPS6 S235/236 and RPS6 was examined via immunoblotting (top). The bottom panel indicates blot intensities relative to control blots (no drug treatment). C, sorafenib-resistant SNU-423 cells were treated with the indicated concentrations of Cl-1040, and relative cell viability was determined 72 h later. Note that Cl-1040 had no significant inhibitory effect on cell growth at 0.1 μ M (indicated by a red arrow). D, sorafenib-resistant SNU-423 cells were treated with the indicated concentrations of AZD8055 in the presence (open circles) or absence (solid circles) of 0.1 μ M Cl-1040, and relative cell viability was determined 72 h later.

AZD8055, than sorafenib-sensitive cell lines (Fig. 3*E*), indicating that sorafenib-resistant cells are dependent for growth on constitutive activation of the mTOR pathway.

Synergy of mTOR and MAPK Inhibitors—Although JHH-1 and SNU-182 cells showed resistance to sorafenib, their ERK phosphorylation was dose-dependently attenuated by sorafenib (Fig. 3C), indicating that the MAPK pathway in these

cells still retained some sensitivity to the inhibition of RAF or other unknown MAPK-pathway kinases. In fact, sorafenib augmented the down-regulation of p-RPS6 S235/236 by everolimus (Fig. 4A). It is noteworthy that the low level of p-RPS6 S235/236 in SNU-182 cells sustained in the presence of 2 μ M everolimus was completely abrogated by the addition of 3 μ M sorafenib.

Fig. 3. **Alternative mTOR signal activation in sorafenib-resistant HCC cells.** *A*, *C*, representative sorafenib-sensitive (HUH-6 and HuH-7) and -resistant (JHH-1, JHH-2, SNU-182, SNU423, and SNU-387) HCC cells were treated with the indicated concentrations of sorafenib for 3 h, and the expression of p-RPS6 S235/236, total RPS6, p-ERK1/2 T202/Y204, total ERK, p-RSK T380, total RSK, p-S6K T389, and total S6K was determined via immunoblotting. *B*, schematic representation of the mTOR and MAPK pathways and their inhibitors. *D*, representative sorafenib-resistant (JHH-1, JHH-2, SNU-182, SNU423, and SNU-387) HCC cells were treated with the indicated concentrations of everolimus for 3 h, and the expression of p-RPS6 S235/236 and total RPS6 was determined via immunoblotting. *E*, distribution of IC₅₀ values of representative sorafenib-sensitive (SNU-449, HUH-6, JHH-7, HuH-7, and SK-Hep1) and -resistant (JHH-1, JHH-2, SNU-182, SNU-423, and SNU-387) HCC cells to AZ8055. Boxes indicate 25th to 75th percentiles.

The phosphorylation of RPS6 S235/236 in JHH-2, SNU-423, and SNU-387 cells was insensitive to sorafenib (Fig. 3C). However, the active MAPK pathway in these cell lines seems to lie downstream of RAF kinases. An MEK inhibitor, CI-1040 (Fig. 3B), enhanced the attenuation of p-RPS6 S235/236 by everolimus in JHH-2 and SNU-423 cells (Fig. 4B), and an RSK inhibitor, SL0101, enhanced the attenuation of p-RPS6 S235/236 by everolimus in JHH-2 and SNU-387 cells (Fig. 4B).

Marked inhibition of p-RPS6 S235/236 by combined blockade of the MAPK pathway downstream of RAF and the mTOR pathway prompted us to examine the effect of this drug combination on HCC cell growth. CI-1040 had no inhibitory effect on the growth of SNU-423 cells at a concentration of $0.1~\mu M$ (Fig. 4C), but it was able to enhance the growthinhibitory effect of AZD8055 (Fig. 4D). Synergy between AZD8055 and CI-1040 was confirmed via Chou-Talalay median dose effect analysis (17) (supplemental Table S8). The combination index values at 50%, 75%, and 90% growth inhibition were 0.783, 0.804, and 0.827, respectively (values of <1 are defined as representative of synergistic effects). These results suggest that HCC patients refractory to sorafenib with a high level of p-RPS6 S235/236 might be treatable with an mTOR inhibitor in combination with drugs that block the MAPK signaling pathway.

To further provide a rational basis for synergistic targeting of the mTOR and MAPK pathways in HCC, we performed unsupervised hierarchical cluster analysis of 23 HCC cell lines based on their phosphorvlation status of signaling components in the mTOR and MAPK pathways listed in supplemental Table S9. Clustering analysis stratified the cell lines into two major groups, A and B (supplemental Fig. S1). In comparison with group A, group B showed higher levels of phosphorylated MAPK signaling components including p-PDGF receptor-β(Thr751), p-Raf-A(Ser299), and phosphorylated signaling modules of the JNK and p38 MAPK pathways (supplemental Fig. S1, C1). Among them, the levels of p-p53 at Ser392, Ser37, and Ser6 differed substantially between groups A (low) and B (high) (supplemental Fig. S1). In addition, cell lines clustered into group A tended to have simultaneous phosphorylation of the mTOR signaling components C2, RPS6(Ser235/236), RPS6(Ser240/244), and eIF4G(Ser1108) (supplemental Fig. S1). With some notable exceptions, sorafenib-insensitive cell lines (high IC_{50} values for sorafenib) and sorafenib-sensitive cell lines (low IC50 values for sorafenib) were clustered into group A and group B, respectively. Although sorafenib-insensitive SNU-387, JHH-1, and KIM-1 cells were classified into the sorafenib-sensitive group B, their phosphorylation levels of mTOR signaling components C2 were higher than those in the other cell lines in group B, suggesting that activation of mTOR signaling might be responsible for the resistance to sorafenib in these cell lines. Some of the sorafenib-insensitive cell lines (e.g. Alexander, JHH-2, and SNU-475 cells) partitioned into subtype Ab (supplemental Fig. S1) were characterized by prominent activation of mTOR signaling components C2 and MAPK signaling components C1. Together, these findings imply that there is a certain population of HCC cells showing up-regulation of both mTOR and MAPK signaling. Such an HCC subtype might respond better to combination treatment with mTOR and MAPK inhibitors. When we compared the phosphorylation status of the mTOR and MAPK signaling nodes in 95 cell lines by means of unsupervised hierarchical clustering, HCC cell lines were significantly clustered together (p = 0.011 by Fisher's exact test) in group A (supplemental Fig. S2), characterized by high levels of phosphorylation of the signaling components C1 and C2, in comparison to cell lines derived from seven other cancer types (supplemental Table S10). The signaling components C1 included previously reported targets of sorafenib such as b-RAF and PDGF receptor- β , as well as the upstream modules of the mTOR pathway (e.g. Akt and PDK). The components C2 comprised RPS6(Ser235/ 236), RPS6(Ser240/244), eIF4G(Ser1108), and the signaling modules of the JNK and p38 MAPK pathways. These observations may reflect the fact that the activation of MAPK signaling by itself, or in combination with mTOR signaling, is a unique feature of HCC.

It is noteworthy that some sorafenib-sensitive and -insensitive cell lines were clustered together into subgroup Aa (supplemental Fig. S1). Although this subgroup was characterized by relatively low levels of both mTOR and MAPK signaling activation, the most sorafenib-sensitive cell line, SNU-449, was classified into this subgroup. It is therefore plausible that some other signaling pathway, in addition to the mTOR and MAPK pathways, may be involved in defining the marked sensitivity of SNU-449 cells to sorafenib.

DISCUSSION

Derangements in the phosphorylation of signaling molecules are hallmarks of cancers, and are often considered as targets of molecular therapies. By profiling the phosphorylation status of multiple signaling components, it is possible to derive important clues for understanding the pathogenesis and classification of cancers. In this study, a high level of p-RPS6 S235/236 was detected in sorafenib-resistant HCC cells. Consistent with this in vitro observation, such high expression of p-RPS6 S235/236 was detected in pretreatment biopsy specimens from HCC patients who had shown early radiographically evident disease progression after starting sorafenib therapy. The number of patient samples analyzed in this study was small and insufficient for providing conclusive evidence, but the present findings warrant future clinical studies to evaluate the significance of p-RPS6 S235/236 as a predictor of response to sorafenib. In order to ensure accurate validation of the utility of p-RPS6 S235/236 as a predictor in future studies, standardized guidelines of immunohistochemistry for detecting p-RPS6 (Ser235/236) need to be developed, including tissue preparation, fixation, staining methods, scoring system, and the definition of a "positive" result.

p-RPS6 has been used as a molecular surrogate for mTOR activation. Villanueva et al. (22) assessed 314 surgical specimens of HCC immunohistochemically using an anti-p-RPS6 S240/244 antibody. They detected p-RPS6 S240/244 in half of the cases examined, and positive staining was correlated with HCC recurrence (22). Although antibodies against p-RPS6 S235/236 and p-RPS6 S240/244 have been used equivalently in many studies to evaluate mTOR activation (22, 23), the phosphorylation of these serine residues was found to be differentially regulated (Fig. 2A). An earlier study demonstrated persistent phosphorylation of RPS6 S235/236 in cells derived from S6K1-/-/S6K2-/- double-knockout mice, and it was concluded that this paradoxical phosphorylation was caused by MAPK signaling. A later study revealed that RSK (MAPK pathway) predominantly phosphorylated the serine 235 and 236 residues of RPS6, whereas S6K (mTOR pathway) broadly phosphorylated the serine 235, 236, 240, 244, and 247 residues. Therefore, use of an antibody against p-RPS6 S240/244 would seem more appropriate for specific detection of the mTOR pathway activation status (20). In the present study, however, we found that phosphorylation of the serine 235 and 236 residues of RPS6 reflected cross-talk between the mTOR and MAPK pathways (Fig. 3C) and served as a predictive biomarker of sorafenib sensitivity. Although RAF kinases (MAPK pathway) are one of the main molecular types targeted by sorafenib, intervention of active mTOR signaling in the MAPK pathway seems to be one of the molecular mechanisms responsible for the resistance of HCC to sorafenib.

It is therefore conceivable that HCC patients with tumors having high levels of p-RPS6 S235/236 could benefit from inhibition of mTOR signaling. We found that mTOR inhibitors showed greater antitumor activity against sorafenib-resisntant HCC cells (Fig. 3E). A recent phase I/II study of everolimus given daily as a single agent in patients with advanced HCC showed that the drug was well tolerated and exerted preliminary antitumor activity in some patients (24). A phase III EVOLVE-1 randomized trial is now ongoing to evaluate the efficacy of everolimus in HCC patients whose disease progressed during or after sorafenib treatment or who were intolerant to sorafenib (25). This clinical trial is designed to reveal the efficacy of mTOR pathway inhibition for control of sorafenib-resistant HCC and is expected to clarify the significance of our present findings.

Clustering analysis of RPPA data revealed that 6 out of 23 HCC cell lines (Alexander, JHH-2, SNU-475, Huh-7, KIM-1, and JHH-1) had prominent activation of both the MAPK and mTOR pathways, indicating a possible subset of HCC patients who might benefit from a combination of MAPK and mTOR inhibitors (supplemental Fig. S1). We also found synergy between MAPK and mTOR pathway inhibitors in sorafenib-resistant cell lines (Fig. 4). However, there is a need for caution before this can be applied clinically. Activation of MAPK signaling occurred at various levels of RAF/MEK/ERK/

RSK in sorafenib-resistant cells (Fig. 3C). In addition, clustering analysis showed activation of two other major MAPK pathways, the Jun N-terminal kinase (JNK) and p38 MAPK pathways, in more than half the HCC cell lines (supplemental Fig. S1). Cross-talk among three major MAPK pathways (RAF/MEK/ERK, JNK, and p38MAPK) has been reported previously (26). Together, these findings suggest that careful assessment is vital when selecting an appropriate MAPK inhibitor for each individual HCC patient. Despite extensive sequencing of kinase genes, we were unable to identify any alterations in the pathway that might be responsible, indicating the need to expedite pharmacoproteomics for therapy personalization.

The present study highlighted the potential power of the RPPA platform for pathway profiling. We have provided proofof-principle support for the utility of the highly sensitive, highthroughput RPPA platform by identifying a practical biomarker with potential clinical applicability. In this study, we used only well-characterized antibodies with high specificity. The Human Antibody Initiative is an ongoing project to raise at least one monospecific antibody against all >20,000 proteins encoded by the human genome (27). It is anticipated that the completion of this project will greatly accelerate the capability of RPPA. The majority of current molecular targeting drugs are designed to target a particular signaling pathway (28). Precise determination of signaling pathways that are activated in individual patients seems to be essential for obtaining maximum benefit from any given treatment. RPPA requires only a minuscule specimen quantity (e.g. less than 1 ng of protein per array) and is applicable even to small biopsy samples. The potential clinical utility of RPPA for decisionmaking and monitoring of cancer therapeutics is thus enormous.

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Supplementary Figure Legends

Figure S1. Unsupervised hierarchical clustering of the 23 HCC cell lines (horizontal axis) and phosphorylation status of the mTOR and MAPK signaling nodes (vertical axis). Red, white and blue colors indicate high, intermediate and low levels of signaling node phophorylation. Light green and yellow represent signaling components of the MAPK pathway and mTOR pathway, respectively. Signaling components of the MAPK and mTOR pathways highly related to HCC cell line subtypes are labeled C1 and C2, respectively. Clustering revealed formation of 2 major groups, A and B, in the cell lines. The 10 cell lines that showed the highest IC50 for sorafenib are highlighted in orange.

Figure S2. Unsupervised hierarchical clustering of the 95 HCC cell lines (horizontal axis) and phosphorylation status of the mTOR and MAPK signaling nodes (vertical axis). Red, white and blue colors indicate high, intermediate and low levels of signaling node phophorylation. Light green and yellow represent signaling components of the MAPK pathway and mTOR pathway, respectively. Signaling components related to HCC cell line subtypes are labeled C1 and C2, respectively. Clustering revealed formation of 2 major groups, A and B, in the cell lines. The 23 HCC cell lines are highlighted in purple.

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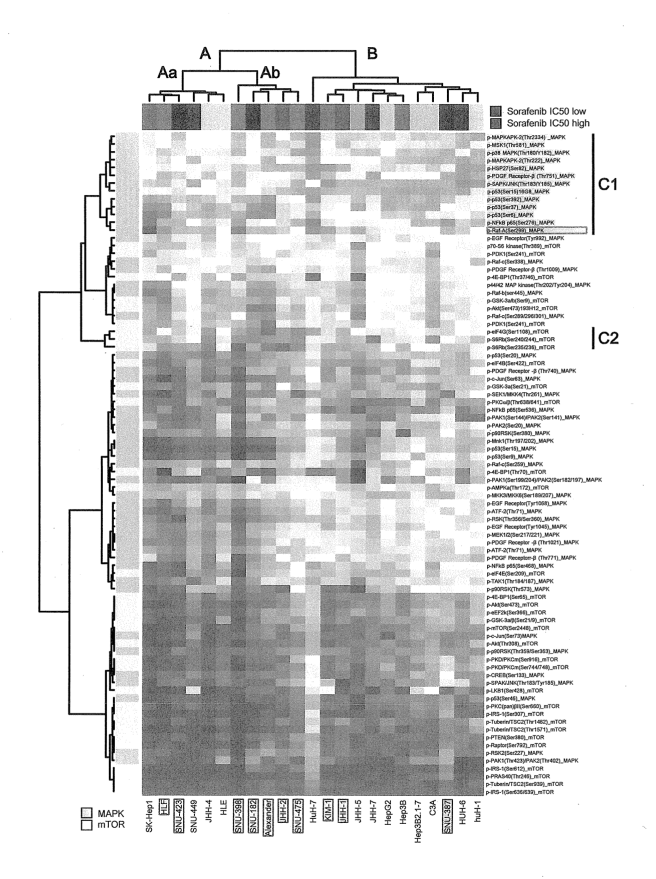
Supplementary Table S6. Kinase gene alterations in HCC cells.

Supplementary Table S7. mTOR and MAPK pathway gene alterations in HCC cells.

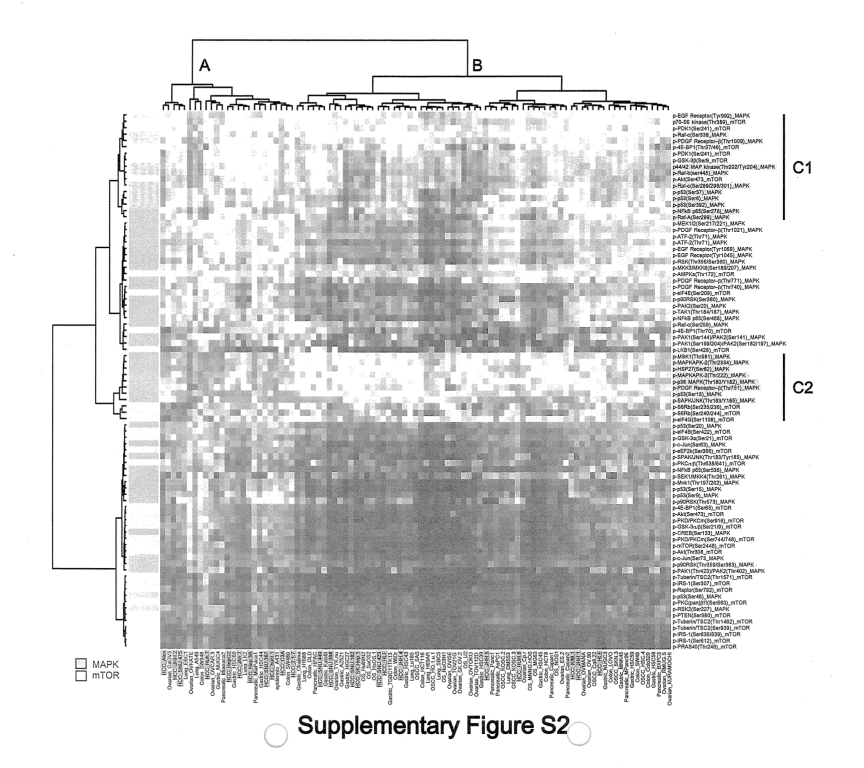
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Supplementary Table S10. Fisher's exact test: Association of cancer types with activation status of MAPK and mTOR signaling.



Supplementary Figure S1.



Supplementary Table S1. List of cell lines used for RPPA construction

#	Cell line	Source	#	Cell line	Source
Iepato	ocellular carcinoma (HCC	C): 23 cell lines	Colon ca	ncer: 9 cell lines	
1	HepG2	JCRB-HSRRB	1	DLD1	JCRB-HSRRB
2	HLE	JCRB-HSRRB	2	HCT-116	ATCC
3	huH-1	JCRB-HSRRB	3	WiDr	JCRB-HSRRB
4	HUH-6 clone 5	JCRB-HSRRB	4	SW1116	ATCC
5	HuH-7	JCRB-HSRRB	5	SW48	ATCC
6	ЈНН-7	JCRB-HSRRB	6	SW948	ATCC
7	HLF	JCRB-HSRRB	7	SW480	ATCC
8	C3A(HepG2/C3A)	ATCC	8	COLO-320	RBCCB
9	Hep 3B2.1-7	ATCC	9	Lovo	JCRB-HSRRB
10	ЈНН-1	JCRB-HSRRB	Pancreat	ic cancer: 9 cell lines	
11	ЈНН-2	JCRB-HSRRB	1	MIAPaCa-1	ATCC
12	ЈНН-4	JCRB-HSRRB	2	MIAPaCa-2	ATCC
13	ЈНН-5	JCRB-HSRRB	3	MPanc-96	ATCC
14	Alexander	ATCC	4	PANC-1	ATCC
15	Hep-3B	JCRB-HSRRB	5	AsPC-1	ATCC
16	KIM-1	JCRB-HSRRB	. 6	BxPC-3	ATCC
17	SK-Hep-1	ATCC	7	Capan-1	ATCC
18	SNU-182	ATCC	8	Capan-2	ATCC
19	SNU-387	ATCC	9	HPAC	ATCC
20	SNU-398	ATCC	Lung can	icer: 8 cell lines	
21	SNU-423	ATCC	1	A549	JCRB-HSRRB
22	SNU-449	ATCC	2	EBC1	JCRB-HSRRB
23	SNU-475	ATCC -	3	LK-2	RBCCB
varia	n cancer: 16 cell lines		4	Lu-165	RBCCB
1	CaOV-3	ATCC	. 5	NCI-H1688	ATCC
2	TYK-nu	JCRB-HSRRB	6	H69AR	ATCC
3	TYK-nu.CP-r	JCRB-HSRRB	7	SBC5	JCRB-HSRRB
4	RMUG-S	JCRB-HSRRB	8	DMS53	ATCC
5	ES-2	ATCC	Osteosar	coma: 8 cell lines	
6	SK-OV-3	ATCC	1	U-2	ATCC
7	OV-90	ATCC	2	SaOS2	ATCC
8	TOV-112D	ATCC	3	NOS1	RBCCB
9	TOV-21G	ATCC	4	HuO 9N2	RBCCB
10	KURAMOCHI	JCRB-HSRRB	5	HuO-3N1	RBCCB
11	OVCAR-3	ATCC	6	MNNG/HOS	ATCC
12	OVISE	JCRB-HSRRB	7	MG-63	RBCCB
13	OVKATE	JCRB-HSRRB	8	Hs-OS-1	RBCCB
14	OVMANA	JCRB-HSRRB	Oral can	with the end togeth swift with a least part of the control	
15	OVSAHO	JCRB-HSRRB	1	SAS	JCRB-HSRRB
16	OVTOKO	JCRB-HSRRB	2	Ca9-22	RBCCB
	c cancer: 15 cell lines		3	Ho-1-u-1	JCRB-HSRRB
1	HSC-39	Suppl. Ref [1]	4	HSC-4	RBCCB
2	HSC-43	Suppl. Ref [1]	5	KOSC-2	JCRB-HSRRB
3	HSC-44	Suppl. Ref [1]	6	KOSC-3	JCRB-HSRRB
4	HSC-45	Suppl. Ref [1]	7	SKN-3	JCRB-HSRRB
5	HSC-58	Suppl. Ref [1]		oid carcinoma: 1 cell	
6	HSC-60	Suppl. Ref [1]	1	A431	RBCCB
7	HSC-59	Suppl. Ref [1]	nonement summer services		
8	MKN-45	JCRB-HSRRB	Abbrevia	tions: JCRB-HSRPR	Japanese Collection of Res
9	Okajima	Suppl. Ref [2]			search Resources Bank (Os
10	AZ521	RBCCB			Culture Collection (Manass
11	Kato-III	RBCCB			rce Center Cell Bank (Tsu
12	NUGC-3	JCRB-HSRRB	Japan).	,	(100
12	NILICC 4	ICDD HEDDD	_F /,		

13

14

15

NUGC-4

HGC27

TGBC11TKB

JCRB-HSRRB

RBCCB

RBCCB

[†]EGF treated A431 cells were included as a control.

SUPPLEMENTARY REFERENCES

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Supplementary Table S2. List of 180 phosphorylation site-specific antibodies

Supplier	Cat.#	Ab name	Pathway	Dilution	Ab type
CST	9570	p-eNOS (Ser1177) C9C3	Angiogenesis	1:1000	Ra mAb
CST	9296	p-Bad(Ser112)	Apoptosis	1:2000	Mo mAb
CST	5286	p-Bad(Ser136) 185D10	Apoptosis	1:500	RamAb
CST	9297	p-Bad(Ser155)	Apoptosis	1:1000	Ra pAb
CST	2827	p-Bcl-2(Ser70) 5H2	Apoptosis	1:1000	Ra mAb
CST	2875	p-Bcl-2 (Thr56) Antibody (Human Specific)	Apoptosis	1:1000	Ra pAb
CST	4585	p-Bim (Ser69) D7E11	Apoptosis	1:1000	Ra mAb
CST	3079	p-Aurora A (Thr288) C39D8	Cell cycle	1:1000	Ra mAb
CST	2914	p-Aurora A (Thr288)/Aurora B (Thr232)/Aurora C (Thr198) D13A11	Cell cycle	1:2000	Ra mAb
CST	3300	p-Cyclin D1 (Thr286) D29B3	Cell cycle	1:1000	Ra mAb
CST	9307	p-Rb (Ser780)	Cell cycle	1:1000	Ra pAb
CST	9301	p-Rb (Ser795)	Cell cycle	1:1000	Ra pAb
CST	9308	p-Rb (Ser807/811)	Cell cycle	1:1000	Ra pAb
CST	4910	p-Wee1 (Ser642) D47G5	Cell Cycle	1:1000	Ra mAb
CST	3843	p-AP2M1	Cytoskelton	1:1000	RapAb
CST	3313	p-Cofilin(Ser3) 77G2	Cytoskelton	1:1000	RamAb
CST	3149	p-Ezrin(Thr567)/Radixin(Thr564)/Moesin(thr558) 41A3	Cytoskelton	1:1000	RamAb
CST	3040	p-MYPT(Ser507)	Cytoskelton	1:1000	RapAb
CST	3048	p-MYPT(Ser668)	Cytoskelton	1:1000	RapAb
CST	4563	p-MYPT(Thr853)	Cytoskelton	1:1000	RapAb
CST	3111	p-VASP(Ser157)	Cytoskelton	1:1000	Ra pAb
CST	3114		Cytoskelton	1:1000	Ra pAb
2002/02/2009/04/04/02/20175/04/		p-VASP(Scr239)		1:1000	Mo mAb
CST	4526	p-ATM(Ser1981) 10H11.E12	DNA damage		
CST	2853	p-ATR(Ser428)	DNA damage	1:1000	Ra pAb
CST	9373	P-BAP1 (Ser592)	DNA damage	1:1000	Ra pAb
CST	9009	p-BRCA1(Ser1524)	DNA damage	1:1000	Ra pAb
CST	2349	p-Chk1 (Ser296)	DNA damage	1:1000	Ra pAb
CST	2344	p-Chk1 (Ser317)	DNA damage	1:1000	Ra pAb
CST	2341	p-Chk1 (Ser345)	DNA damage	1:1000	Ra pAb
CST	2348	p-Chk1 (Ser345) 133D3	DNA damage	1:1000	Ra mAb
CST	2666	p-Chk2 (Ser19)	DNA damage	1:1000	Ra pAb
CST	2665	p-Chk2 (Ser33/35)	DNA damage	1:1000	Ra pAb
CST	2668	p-Chk2 (Thr387)	DNA damage	1:1000	Ra pAb
CST	2661	p-Chk2 (Thr68)	DNA damage	1:1000	Ra pAb
CST	9284	p-p53(Ser15)	DNA damage	1:1000	Ra pAb
CST	9286	p-p53(Ser15) 16G8	DNA damage	1:1000	Mo mAb
CST	9287	p-p53(Ser20)	DNA damage	1:2000	Ra pAb
CST	9289	p-p53(Ser37)	DNA damage	1:1000	Ra pAb
CST	9281	p-p53(Ser392)	DNA damage	1:1000	Ra pAb
CST	2521	p-p53(Ser46)	DNA damage	1:750	Ra pAb
CST	9285	p-p53(Ser6)	DNA damage	1:1000	Ra pAb
CST	9288	p-p53(Ser9)	DNA damage	1:1000	Ra pAb
CST	9461	p-FoxO1 (Ser256)	Forkhead	1:750	Ra pAb
CST	2487	p-FoxO1 (Ser319)/FoxO4(Ser262)	Forkhead	1:1000	Ra pAb
CST	9464	p-FoxO1 (Thr24)/FoxO3a (Thr32)	Forkhead	1:750	Ra pAb
CST	2599	p-FoxO1 (Thr24)/FoxO3a (Thr32)/FoxO4 (Thr28) 4G6	Forkhead	1:1000	Ra mAb
CST	9466	p-FoxO3a (Ser253)	Forkhead	1:1000	Ra pAb
CST	9465	p-FoxO3a (Ser318/321)	Forkhead	1:750	Ra pAb
CST	3661	p-Acetyl-CoA carboxylase(Ser79)	Glucose metabolism	1:1000	Ra pAb
CST	2535	p-AMPKa(Thr172)	Glucose metabolism	1:1000	Ra pAb
CST	4181	p-AMPKb(Ser108)	Glucose metabolism	1:1000	RapAb
CST	3482	p-LKB1 (Ser428) C67A3	Glucose metabolism	1:1000	Ra mAb
CST	2381	p-IRS-1(Ser307)	Glucose metabolism	1:1000	RapAb
CST	3203	p-IRS-1(Ser612) C15H5	Glucose metabolism	1:1000	Ra mAb
		-			Ra pAb
CST	2388	p-IRS-1(Ser636/639)	Glucose metabolism	1:1000	

CST	3827	p-PKM2 (Tyr105)	Glucose metabolism	1:1000	Ra pAb
CST	3776	p-Jak2 (Tyr1007/1008) C80C3	JAK/STAT	1:1000	RamAb
CST	9171	p-Stat 1(Y701)	JAK/STAT	1:1000	Ra pAb
CST	4441	p-Stat 2(Y690)	JAK/STAT	1:1000	Ra pAb
CST	9134	p-Stat 3(Ser727)	JAK/STAT	1:1000	RapAb
CST	9145	p-Stat 3(Y705) D3A7	JAK/STAT	1:3000	RamAb
CST	9359	p-Stat 5(Y694) C11C5	JAK/STAT	1:1000	RamAb
CST	9361	p-Stat 6(Y641)	JAK/STAT	1:1000	RapAb
CST	9321	p-Tyk2 (Tyr1054/1055)	JAK/STAT	1:1000	RapAb
CST	2577	p-Histone H2A.X(Ser139)	Mitosis marker	1:1000	Ra pAb
CST	9701	p-Histone H3(Ser10)	Mitosis marker	1:1000	Ra pAb
CST	9713	p-Histone H3(Ser28)	Mitosis marker	1:1000	Ra pAb
CST	9764	p-Histone H3(Thr11)	Mitosis marker	1:1000	Ra pAb
CST	9714	p-Histone H3(Thr3)	Mitosis marker	1:1000	Ra pAb
CST	2681	p-IKKα (Ser180)/IKKβ (Ser181)	NFkB	1:1000	Ra pAb
CST	9246	p-IκB-α (Ser32/36) 5A5	NFkB	1:1000	Mo mAb
CST	3037	p-NFkB p65(Ser276)	NFkB	1:1000	Ra pAb
CST	3039	p-NFkB p65(Ser468)	NFkB	1:1000	Ra pAb
CST	3033	p-NFkB p65(Ser536)93H1	NFkB	1:1000	Ra mAb
CST	5112	p-ATF-2(Thr71)11G2	p38 MAPK	1:1000	Ra mAb
CST	2401	p-HSP27(Ser82)	p38 MAPK	1:1000	Ra pAb
CST	3316	p-MAPKAPK-2(Thr222) 9A7	p38 MAPK	1:1000	Ra mAb
CST	3007	p-MAPKAPK-2(Thr2334)27B7	p38 MAPK	1:1000	Ra mAb
CST	9236	p-MKK3/MKK6(Ser189/207) 22A8	p38 MAPK	1:1000	Ra mAb
CST	9595	p-MSK1(Thr581)	p38 MAPK	1:1000	Ra pAb
CST	4511	p-p38 MAPK(Thr180/Y182) D3F9	p38 MAPK	1:1000	Ra mAb
CST	9371	p-PKC(pan)βII (Ser660)	Phospholipase	1:1000	Ra pAb
CST	9375	p-PKCα/β(Thr638/641)	Phospholipase	1:1000	Ra pAb
CST	9374	p-PKCΔ(Thr505)	Phospholipase	1:1000	RapAb
CST	9376	p-PKCΔ/θ(Ser643/676)	Phospholipase	1:750	RapAb
CST	9378	p-PKCZ/\(\lambda\)(Thr410/403)	Phospholipase	1:750	RapAb
CST	9377	p-PKC0(Thr538)	Phospholipase	1:750	Ra pAb
CST	3871	p-PLCg(Y1217)	Phospholipase	1:1000	RapAb
CST	3874	p-PLCg(Y759)	Phospholipase	1:1000	RapAb
CST CST	2821 4923	p-PLCg(Y783) Non-p-4E-BP1(Thr46) 87D12	Phospholipase PI3K-AKT-mTOR	1:1000 1:1000	Ra pAb Ra mAb
CST	9451	p-4E-BP1 (Ser65)	PI3K-AKT-mTOR	1:1000	
CST	2855	p-4E-BP1 (3603)	PI3K-AKT-mTOR	1:1000	Ra pAb Ra pAb
CST	9455	p-4E-BP1 (Thr70)	PI3K-AKT-mTOR	1:1000	Ra pAb
CST	9234	p70-S6 kinase(Thr389)	PI3K-AKT-mTOR	1:1000	Ra pAb
CST	4058	Phospho-Akt (Ser473) 193H12	PI3K-AKT-mTOR	1:2000	Ra mAb
CST	2965	p-Akt(Thr308)C31E5	PI3K-AKT-mTOR	1:1000	Ra mAb
CST	3691	p-eEF2k(Ser366)	PI3K-AKT-mTOR	1:1000	Ra pAb
CST	3591	p-elF4B(Ser422)	PI3K-AKT-mTOR	1:1000	Ra pAb
CST	9741	p-elF4E(Ser209)	PI3K-AKT-mTOR	1:1000	Ra pAb
CST	2441	p-elF4G(Ser1108)	PI3K-AKT-mTOR	1:1000	Ra pAb
CST	9316	p-GSK-3a(Ser21)36E9	PI3K-AKT-mTOR	1:1000	Ra mAb
CST	9327	p-GSK-3a/β(Ser21/9)37F11 (GSK-3a preferred)	PI3K-AKT-mTOR	1:1000	Ra mAb
CST	9323	p-GSK-3β(Ser9)5B3	PI3K-AKT-mTOR	1:1000	Ra mAb
CST	2971	p-mTOR(Ser2448)	PI3K-AKT-mTOR	1:1000	Ra pAb
CST	3061	p-PDK1(Ser241)	PI3K-AKT-mTOR	1:1000	Ra pAb
CST	2054	p-PKD/PKCm(Ser744/748)	PI3K-AKT-mTOR	1:1000	Ra pAb
CST	2051	p-PKD/PKCm(Ser916)	PI3K-AKT-mTOR	1:1000	Ra pAb
CST	2997	p-PRAS40(Thr246) C77D7	PI3K-AKT-mTOR	1:1000	Ra mAb
CST	9551	p-PTEN(Ser380)	PI3K-AKT-mTOR	1:1000	Ra pAb
CST	2083	p-Raptor(Ser792)	PI3K-AKT-mTOR	1:1000	Ra pAb
CST	3556	p-RSK2 (Ser227) D53A11	PI3K-AKT-mTOR	1:1000	Ra mAb
CST	4858	p-S6Rb(Ser235/236) D57.2.2e	PI3K-AKT-mTOR	1:2000	Ra mAb
CST	4838	p-S6Rb(Ser240/244)	PI3K-AKT-mTOR	1:1000	Ra pAb

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CST	3615	p-Tuberin/TSC2 (Ser939)	PI3K-AKT-mTOR	1:1000	Ra pAb
CST	3617	p-Tuberin/TSC2 (Thr1462) 5B12	PI3K-AKT-mTOR	1:1000	Ra mAb
CST	3614	p-Tuberin/TSC2 (Thr1571)	PI3K-AKT-mTOR	1:1000	Ra pAb
CST	9101	p44/42 MAP kinase(Thr202/Tyr204)	Ras-Raf	1:1000	Ra pAb
CST CST	9198	p-CREB (Ser133) 87G3	Ras-Raf	1:1000	Ra mAb Ra mAb
CST	9154	p-MEK1/2 (Ser217/221) 41G9	Ras-Raf Ras-Raf	1:1000 1:1000	
CST	2111 9335	p-Mnk1 (Thr197/202)	Ras-Raf	1:1000	Ra pAb Ra mAb
CST	9344	p-p90RSK(Ser380) 9D9 p-p90RSK(Thr359/Ser363)	Ras-Raf	1:1000	Ra pAb
CST	9344		Ras-Raf	1:1000	
CST	2606	p-p90RSK(Thr573) p-PAK1(Ser144)PAK2(Ser141)	Ras-Raf	1:1000	Ra pAb Ra pAb
CST	2605	p-PAK1(Ser199/204)PAK2(Ser182/197)	Ras-Raf	1:1000	Ra pAb
CST	2601	p-PAK1(Thr423)/PAK2(Thr402)	Ras-Raf	1:1000	Ra pAb
CST	2607	p-PAK2(Ser20)	Ras-Raf	1:1000	Ra pAb
CST	4431	p-Raf-A (Ser299)	Ras-Raf	1:1000	RapAb
CST	2696	p-Raf-b(ser445)	Ras-Raf	1:1000	Ra pAb
CST	9421	p-Raf-c (Ser259)	Ras-Raf	1:1000	RapAb
CST	9431	p-Raf-c(Ser289/296/301)	Ras-Raf	1:1000	RapAb
CST	9427	p-Raf-c(Ser338)56A6	Ras-Raf	1:1000	RamAb
CST	9348	p-RSK(Thr356/Ser360)	Ras-Raf	1:1001	RapAb
CST	2102	Non-p-Src(Y416)7G9	RTK/TK	1:1000	Mo mAb
CST	2102	Non-p-Src(Y527)	RTK/TK	1:1000	Ra pAb
CST	3714	pALK (Tyr1282/1283)	RTK/TK	1:1000	Ra pAb
CST	3073	p-c-Kit(Y703)D12E12	RTK/TK	1:1000	Ra mAb
CST	2237	p-EGF Receptor (Tyr1045)	RTK/TK	1:1000	Ra pAb
CST	3777	p-EGF Receptor (Tyr1068) D7A5	RTK/TK	1:1000	Ra mAb
CST	2235	p-EGF Receptor (Tyr992)	RTK/TK	1:1000	RapAb
CST	3474	p-FLT3(Y591) 33G6	RTK/TK	1:1000	Ra mAb
CST	3233	p-Gab1 (Tyr627) C32H2	RTK/TK	1:1000	Ra mAb
CST	3882	p-Gab2 (Tyr452) C33G1	RTK/TK	1:1000	Ra mAb
CST	2243	p-HER2/ErbB2 (Tyr1221/1222) 6B12	RTK/TK	1:1000	Ra mAb
CST	2247	p-HER2/ErbB2 (Tyr1248)	RTK/TK	1:1000	Ra pAb
CST	3790	p-HER4/ErbB4 (Tyr984)	RTK/TK	1:1000	Ra pAb
CST	3021	p-IGF-I Receptor (Tyr1131)/Insulin Receptor (Tyr1146)	RTK/TK	1:1000	Ra pAb
CST	3024	p-IGF-I Receptor (Tyr1135/1136)	RTK/TK	1:1000	Ra pAb
CST	4568	p-IGF-I Receptor (Tyr980)	RTK/TK	1:1000	Ra pAb
CST	3077	p-Met(Thr1234/1235) D26	RTK/TK	1:1000	Ra mAb
CST	3124	p-PDGF Receptor-β(Thr1009) 42F9	RTK/TK	1:1000	Ra mAb
CST	2227	p-PDGF Receptor-β(Thr1021) 6F10	RTK/TK	1:1000	Ra mAb
CST	3168	p-PDGF Receptor-β(Thr740) 32A9	RTK/TK	1:1000	Ra mAb
CST	4549	p-PDGF Receptor-β(Thr751) C63G6	RTK/TK	1:1000	Ra mAb
CST	3173	p-PDGF Receptor-β(Thr771)	RTK/TK	1:1000	Ra mAb
CST	3221	p-Ret (Tyr905)	RTK/TK	1:1000	Ra pAb
Upstate	209	p-SHC(Y239)	RTK/TK	1:1000	Ra pAb
Upstate	206	p-SHC(Y317)	RTK/TK	1:1000	Ra pAb
CST	2007	p-SHIP2(Tyr1135)	RTK/TK	1:1000	Ra pAb
CST	2101	p-Src Family (Y416)	RTK/TK	1:1000	Ra pAb
CST	2105	p-Src(Y527)	RTK/TK	1:1000	Ra pAb
CST	2715	p-Syk (Tyr323)	RTK/TK	1:1000	Ra pAb
CST	2711	p-Syk (Tyr525/526)	RTK/TK	1:1000	Ra pAb
CST	3817	p-VEGF Receptor 2 (Tyr1059) D5A6	RTK/TK	1:1000	Ra mAb
CST	3770	p-VEGF Receptor 2 (Tyr1175) D5B11	RTK/TK	1:1000	Ra mAb
CST	4991	p-VEGF Receptor 2 (Tyr951) 15D2	RTK/TK	1:1000	RamAb
CST	2701	p-Zap-70 (Tyr319)/Syk (Tyr352)	RTK/TK	1:1000	Ra pAb
CST	9221	p-ATF-2(Thr71)	SAPK/JNK	1:1000	RapAb
CST	9261	p-c-Jun (Ser63) II	SAPK/JNK	1:1000	Ra pAb
CST	3270	p-c-Jun (Ser73) D47G9	SAPK/JNK	1:1000	RamAb
CST	4671	p-SAPK/JNK(Thr183/Y185) 98F2	SAPK/JNK	1:1000	RamAb
CST	9151	p-SEK1/MKK4(Thr261)	SAPK/JNK	1:1000	RapAb
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CST	4668	p-SPAK/JNK(Thr183/Tyr185) 81E11	SAPK/JNK	1:1000	Ra mAb
CST	9516	p-Smad1/5(Ser463/465) 41D10	TGF-β	1:1000	Ra mAb
CST	3108	p-Smad2(Ser465/467) 138D4	TGF-β	1:1000	Ra mAb
CST	4508	p-TAK1 (Thr184/187) 90C7	TGF-β	1:1000	Ra mAb
CST	2568	p-LRP6 (Ser1490)	Wnt	1:1000	Ra pAb
CST	9561	p-β-catenin(Ser33/37/Thr41)	Wnt	1:750	Ra pAb
CST	9564	p-β-catenin(Ser45)	Wnt	1:1000	Ra pAb
Sigma	T6557	γ-Tublin	For nomalization	1:2000	Mo mAb

**Abbreviations:** CST, Cell Signaling Technology, Inc. (Danvers, MA); Upstate, Upstate Biotechnology, Inc. (Lake Placid, NY); Sigma, Sigma-Aldrich, Co. (St. Louis, MO); RTK/TK, Rreceptor tyrosine kinase/tyrosine kinase; Mo, mouse; Ra, rabbit; mAb, monoclonal antibody; pAb, polyconal antibody.

Supplementary Table S3.

Correlation between sensitivity to sorafenib and the phosphorylation of 180 signaling nodes

Rank	Antibody	Spearman CC	P value	Rank	Antibody	Spearman CC	P value
1	p-S6Rb( Ser235/236 ) D57.2.2e	0.5791	0.0044	60	p-p53(Ser46)	-0.0504	0.8196
2	p-S6Rb(Ser240/244)	0.5524	0.0070	61	p-Ret (Tyr905)	-0.0534	0.8091
3	p-RSK2 (Ser227) D53A11	0.3370	0.1156	62	p-Stat 1(Y701)	-0.0613	0.7812
4	p-elF4G(Ser1108)	0.3320	0.1213	63	p-MSK1(Thr581)	-0.0632	0.7742
5	p-NFkB p65(Ser536) 93H1	0.3281	0.1260	64	p-Raf-c (Ser259)	-0.0642	0.7708
6	p-IRS-1(Ser612) C15H5	0.3093	0.1502	65	p-β-catenin(Ser33/37/Thr41)	-0.0692	0.7535
7	p-c-Kit(Y703) D12E12	0.2688	0.2133	66	p-ATR(Ser428)	-0.0702	0.7501
8	p-Chk1 (Ser317)	0.2609	0.2274	67	p-Smad2(Ser465/467) 138D4	-0.0731	0.7398
9	p-Tuberin/TSC2 (Ser939)	0.2470	0.2537	68	p-Tuberin/TSC2 (Thr1571)	-0.0800	0.7160
10	p-Bad(Ser136)185D10	0.2322	0.2839	69	p-Rb (Ser807/811)	-0.0860	0.6958
11	p-90RSK(Ser380) 9D9	0.2292	0.2902	70	p-PRAS40(Thr246) C77D7	-0.0870	0.6925
12	p-Bim (Ser69) D7E11	0.2233	0.3031	71	P-BAP1 (Ser592)	-0.0968	0.6593
13	p-PAK1(Thr423)/PAK2(Thr402)	0.2134	0.3254	72	p-Met(Thr1234/1235) D26	-0.1077	0.6236
14	Non-p-Src(Y527)	0.1937	0.3730	73	p-Stat 5(Y694) C11C5	-0.1087	0.6203
15	p-Cyclin D1 (Thr286) D29B3	0.1897	0.3829	74	p-Stat 3(Ser727)	-0.1097	0.6171
16	p-Chk1 (Ser345)	0.1729	0.4270	75	p-p90RSK(Thr359/Ser363)	-0.1126	0.6076
17	p-PAK1(Ser199/204)PAK2(Ser182/197)	0.1670	0.4432	76	p-Rb (Ser795)	-0.1166	0.5949
18	p-PKCZ/λ(Thr410/403)	0.1611	0.4597	77	p-Acetyl-CoA carboxylase(Ser79)	-0.1206	0.5824
19	p-PTEN(Ser380)	0.1512	0.4879	78	p-Bcl-2(Thr56)	-0.1245	0.5699
20	p-PAK1(Ser144)PAK2(Ser141)	0.1344	0.5379	79	p-LRP6 (Ser1490)	-0.1255	0.5669
21	p-ATM(Ser1981) 10H11.E12	0.1324	0.5440	80	p-Bad(Ser155)	-0.1275	0.5607
22	p-PKCα/β(Thr638/641)	0.1324	0.5933	81	p-MEK1/2 (Ser217/221) 41G9	-0.1314	0.5485
	• • • • • • • • • • • • • • • • • • • •					-0.1364	0.5334
.23	p-HER2/ErbB2 (Tyr1221/1222) 6B12	0.1136	0.6028	82	p-MYPT(Ser668)		
24	p-PKC0(Thr538)	0.1136	0.6028	83	p-Histone H3(Ser28)	-0.1374	0.5304
25	p-AMPKa(Thr172)	0.1117	0.6119	84	p-p53(Ser20)	-0.1443	0.5097
26	p-Raptor(Ser792)	0.1107	0.6123	85	p-Src(Y527)	-0.1443	0.5097
27	p-HER2/ErbB2 (Tyr1248)	0.0988	0.6511	86	p-Src Family (Y416)	-0.1443	0.5097
28	p-Chk2 (Ser19)	0.0879	0.6874	87	p-eEF2k(Ser366)	-0.1462	0.5039
29	p-eNOS (Ser1177) C9C3	0.0800	0.7143	88	p-PKD/PKCm(Ser744/748)	-0.1462	0.5039
30	p-Ezrin(Thr567)/Radixin(Thr564)/Moesin(thr558)	0.0771	0.7245	89	p-EGF Receptor (Tyr1045)	-0.1482	0.4980
31	p-FLT3(Y591) 33G6	0.0731	0.7381	90	p-Gab2 (Tyr452) C33G1	-0.1571	0.4723
32	p-Wee1 (Ser642) D47G5	0.0573	0.7933	91	p-p38 MAPK(Thr180/Y182 ) D3F9	-0.1581	0.4695
33	p-EGF Receptor (Tyr992)	0.0553	0.8003	92	p-elF4B(Ser422)	-0.1611	0.4611
34	p-IRS-1(Ser636/639)	0.0455	0.8354	93	p-PKD/PKCm(Ser916)	-0.1611	0.4611
35	p-Tuberin/TSC2 (Thr1462) 5B12	0.0336	0.8780	94	p-PKCΔ(Thr505)	-0.1640	0.4528
36	p-elF4E(Ser209)	0.0296	0.8923	95	p-BRCA1(Ser1524)	-0.1650	0.4500
37	p-GSK-3a(Ser21) 36E9	0.0267	0.9030	96	p-Chk2 (Thr68)	-0.1680	0.4418
38	p-PAK2(Ser20)	0.0257	0.9066	97	p-LKB1 (Ser428) (C67A3)	-0.1680	0.4418
39	p-Smad1/5(Ser463/465) 41D10	0.0227	0.9173	98	p-FoxO3a (Ser318/321)	-0.1700	0.4364
40	p-PDGF Receptor-β(Thr740)	0.0198	0.9281	99	p-p53(Ser9)	-0.1709	0.4337
41	p-MKK3/MKK6(Ser189/207) 22A8	0.0168	0.9388	100	p-4E-BP1 (Thr70)	-0.1779	0.4151
42	p-PKCΔ/θ(Ser643/676)	0.0148	0.9460	101	p-IGF-I Receptor (Tyr1135/1136)	-0.1789	0.4124
43	p-Raf-A (Ser299)	0.0030	0.9892	102	p-CREB (Ser133) 87G3	-0.1818	0.4046
44	p-HSP27(Ser82)	0.0020	0.9928	103	p-PLCg(Y759)	-0.1828	0.4020
	p-Raf-c(Ser338) 56A6					-0.1877	0.3892
45 46	, , ,	0.0010	0.9964	104	p-IRS-1(Ser307)		
46	p-Chk1 (Ser296)	-0.0079	0.9730	105	p-mTOR(Ser2448)	-0.1927	0.3767
47	p-AP2M1	-0.0089	0.9694	106	p-Mnk1 (Thr197/202)	-0.1927	0.3767
48	p-4E-BP1 (Thr37/46)	-0.0109	0.9622	107	p-VASP(Ser157)	-0.1996	0.3595
49	p-Aurora A (T288)/Aurora B (T232)/Aurora C (T198)		0.9586	108	p-Chk2 (Ser33/35)	-0.2016	0.3547
50	p-PLCg(Y783)	-0.0178	0.9370	109	p-lκB-α (Ser32/36) 5A5	-0.2036	0.3499
51	p-SPAK/JNK(Thr183/Tyr185) 81E11	-0.0287	0.8976	110	p-Zap-70 (Tyr319)/Syk (Tyr352)	-0.2036	0.3499
52	p-TAK1 (Thr184/187) 90C7	-0.0356	0.8727	111	p-Chk2 (Thr387)	-0.2065	0.3428
53	p-IGF-I R(Tyr1131)/Insulin R (Tyr1146)	-0.0366	0.8691	112	Non-p-Src(Y416) 7G9	-0.2085	0.3381
54	p-Cofilin(Ser3) 77G2	-0.0425	0.8478	113	p-Tyk2 (Tyr1054/1055)	-0.2125	0.3289
55	p-Rb (Ser780)	-0.0435	0.8443	114	p-PKC(pan)βII (Ser660)	-0.2144	0.3243
56	p-FoxO1 (Ser319)/FoxO4(Ser262)	-0.0445	0.8407	115	p-VEGF Receptor 2 (Tyr1175)	-0.2164	0.3198
57	p-VASP(Ser239)	-0.0494	0.8231	116	p-PDGF Receptor-β(Thr751)	-0.2174	0.3175
			0.8231	117	p-PKM2 (Tyr105)	-0.2174	0.3175
58	p-GSK-3a/β(Ser21/9) 37F11	-0.0494	0.0231	117	p-1 KWZ (191100)	-0.2114	0.0170