www.nature.com/emm

ORIGINAL ARTICLE

Aldehyde dehydrogenase is used by cancer cells for energy metabolism

Joon Hee Kang^{1,8}, Seon-Hyeong Lee^{1,8}, Dongwan Hong², Jae-Seon Lee¹, Hee-Sung Ahn^{3,4}, Ju-Hyun Ahn^{5,6}, Tae Wha Seong¹, Chang-Hun Lee¹, Hyonchol Jang¹, Kyeong Man Hong⁷, Cheolju Lee^{3,4}, Jae-Ho Lee^{5,6} and Soo-Youl Kim¹

We found that non-small-cell lung cancer (NSCLC) cells express high levels of multiple aldehyde dehydrogenase (ALDH) isoforms via an informatics analysis of metabolic enzymes in NSCLC and immunohistochemical staining of NSCLC clinical tumor samples. Using a multiple reaction-monitoring mass spectrometry analysis, we found that multiple ALDH isozymes were generally abundant in NSCLC cells compared with their levels in normal IMR-90 human lung cells. As a result of the catalytic reaction mediated by ALDH, NADH is produced as a by-product from the conversion of aldehyde to carboxylic acid. We hypothesized that the NADH produced by ALDH may be a reliable energy source for ATP production in NSCLC. This study revealed that NADH production by ALDH contributes significantly to ATP production in NSCLC. Furthermore, gossypol, a pan-ALDH inhibitor, markedly reduced the level of ATP. Gossypol combined with phenformin synergistically reduced the ATP levels, which efficiently induced cell death following cell cycle arrest.

Experimental & Molecular Medicine (2016) 48, e272; doi:10.1038/emm.2016.103; published online 25 November 2016

INTRODUCTION

Although tyrosine kinase inhibitor therapy achieves higher objective response rates and longer remissions in lung cancer, relapses occur after 10–14 months owing to drug resistance. 1,2 Therefore, understanding the mechanisms that underlie the biology of non-small-cell lung cancer (NSCLC) cells may provide a chance to improve therapeutic approaches by targeting metabolic enzymes involved in this drug resistance. Therefore, we explored changes in the expression of metabolic enzymes in NSCLC using a bioinformatics analysis. Our analysis revealed that aldehyde dehydrogenase isoforms (EC 1.2.1.3, ALDH) were upregulated in patients with lung adenocarcinoma (LUAD). Several reports have demonstrated an association between ALDH and clinical factors associated with cancer stemness in NSCLC. Survival analyses have revealed that ALDH1L1-positive patients have shorter overall survival rates than ALDH1L1-negative patients.^{3,4} Furthermore, a xenograft model has been used to show that ALDH1A1-positive NSCLC cells are 100 times more tumorigenic than ALDH1A1-negative NSCLC cells.³ Although ALDH1A1- and ALDH3A1-positive tumors are considered to be malignant, the specific role of ALDH in the growth and survival of NSCLC remains unclear.

As a result of the reaction catalyzed by ALDH, NADH is produced from the conversion of aldehyde to carboxylic acid. Here, we investigated the expression of multiple ALDH isoforms in NSCLC, and we tested the role of ALDH in NSCLC to determine whether the NADH produced by ALDH may be a reliable energy source for ATP production in NSCLC. Furthermore, we assessed the effect of gossypol, a pan-ALDH inhibitor, on ATP production and cell death in NSCLC.

MATERIALS AND METHODS

mRNA profiling of LUAD metabolic targets

Gene expression data from RNA sequencing of patients with LUAD were collected from The Cancer Genome Atlas.⁵ Paired sequencing data consisting of matched cancerous and normal tissues were

¹Cancer Cell and Molecular Biology Branch, National Cancer Center, Research Institute, Goyang, Gyeonggi-do, Republic of Korea; ²Cancer Immunology Branch, National Cancer Center, Research Institute, Goyang, Gyeonggi-do, Republic of Korea; ³Center for Theragnosis, Biomedical Research Institute, Korea Institute of Science and Technology, Seoul, Republic of Korea; ⁴Department of Biological Chemistry, University of Science and Technology, Daejeon, Republic of Korea; ⁵Department of Biochemistry and Molecular Biology, Ajou University School of Medicine, Genomic Instability Research Center, Suwon, Korea; ⁶Department of Biomedical Sciences, The Graduate School, Ajou University, Suwon, Korea and ⁷Omics Core Laboratory, National Cancer Center, Research Institute, Goyang, Republic of Korea

⁸These authors contributed equally to this work.

Correspondence: Dr S-Y Kim, Cancer Cell and Molecular Biology Branch, National Cancer Center, Research Institute, 323 Ilsan-ro, Ilsandong-gu, Goyang, Gyeonggi-do 410-769, Republic of Korea.

E-mail: kimsooyoul@gmail.com

Received 20 April 2016; revised 16 June 2016; accepted 29 June 2016

normalized based on RNA sequencing expression estimates via expectation maximization using a custom-made script. To explore metabolic targets associated with cancer, we selected metabolic genes based on the Kyoto Encyclopedia of Genes and Genomes database. Finally, we performed a hierarchical clustering analysis on these genes using the Cluster 3.0 software program (http://bonsai.hgc.jp/~mde-hoon/software/cluster) and visualized the clustered genes using Java Treeview (https://www.princeton.edu/~abarysh/treeview). Student's *t*-tests were used to determine statistical significance between the differentially expressed genes among the clusters. A *P*-value < 0.05 was considered statistically significant.

Cell culture

NCI 60 cell lines, including all NSCLC cell lines, were obtained from the US National Cancer Institute (NCI; MTA no. 2702-09). Cells were incubated at 37 °C and maintained at 5% CO2. H23, H226, IMR-90 (normal lung fibroblasts, ATCC CCL-186) and lung primary (Primary Small Airway Epithelial Cells; Normal, Human, ATCC PCS-301-010) cell lines were obtained from American type culture collection (ATCC, Manassas, VA, USA). NSCLC cells were grown in RPMI 1640 medium (SH30027.01, HyClone, Logan, UT, USA) containing 10% fetal bovine serum (SH30070.03HI, HyClone), penicillin and streptomycin. IMR-90 cells were grown in DMEM/HIGH GLUCOSE medium (SH30243.01, HyClone) containing 10% FBS. Lung primary cells were grown in airway epithelial cell basal medium (PCS-300-030, ATCC) using a bronchial epithelial cell growth kit (PCS-300-040, ATCC). For ALDH overexpression, p3x FLAG-CMV-ALDH isoform constructs individually expressing 1A1, 1A3, 1B1, 2, 3A1, 3A2, 3B1, 4A1, 5A1, 6A1, 7A1, 9A1, 1L1, 1L2 or 18A1 were produced by Cosmogenetech (Seoul, Korea). Each ALDH isoform complementary DNA sequence was obtained from the National Center for Biotechnology Information (NCBI). The ALDH plasmids were transfected into cells using Lipofectamine 3000 (Thermo Fisher Scientific Inc., Fremont, CA, USA) according to the manufacturer's instructions.

Immunoblotting

Cells were harvested and washed in phosphate-buffered saline (PBS) and then lysed in a buffer containing 20 mm Tris-HCl (pH 7.4), 150 mm NaCl, 1% (v/v) Triton X-100, 1 mm EDTA and protease inhibitors before undergoing an immunoblot analysis. The anti-β-actin antibody was purchased from Sigma-Aldrich (A5441, St Louis, MO, USA) and anti-ALDH1A3 (ab129815), anti-ALDH2 (ab54828), anti-ALDH3A2 (ab113111), anti-ALDH3B1 (ab93223) and anti-ALDH3B2 (ab96170) antibodies were purchased from Abcam (Cambridge, UK). Anti-ALDH3A1 (sc-137168), anti-cyclin B1 (sc-245) and anti-α-tubulin (sc-23948) antibodies were purchased from Santa Cruz Biotechnology (Santa Cruz, CA, USA). The anti-pY15-Cdk1 antibody was purchased from Cell Signaling Technology (Dallas, TX, USA, #9111S).

Analysis of ALDH isozymes via multiple reactionmonitoring mass spectrometry

Sample preparation. One primary lung and eight lung cancer cell lines (A549, EKVK, H322M, H23, H460, Hop62, Hop92 and IMR-90) were prepared. The harvested cells were resuspended in 400 μ l of chilled lysis buffer containing 8 M urea, 50 mm Tris (pH 8.0), 64 mm sodium pyrophosphate, 1 mm sodium orthovanadate, 30 mm NaCl, 1× protease inhibitor cocktail and 1× phosphatase inhibitor cocktail. The cells were sonicated three times with 1-s pulses separated by 19 s of cooling at 4 °C. The lysates were centrifuged at 13 400 g for 10 min

at 4 °C to remove cellular debris. Supernatants were collected, and the total protein amount was quantified using the BCA assay (Thermo, Waltham, MA, USA). The protein concentration was adjusted to 0.6 mg ml⁻¹ using the lysis buffer.

Each cell lysate (30 µg in 50 µl) was mixed with 15 µl of 50 mm Tris (pH 8.0) and 10 µl of 50 mm tris-(2-carboxyethyl)-phosphine and then incubated at 25 °C for 1 h. To the reaction mixture, 20 µl of 100 mm iodoacetamide was added, and mixtures were incubated for an additional 1 h in darkness. The samples were diluted fourfold with 50 mm Tris (pH 8.0) to reduce the urea concentration to less than 1 M. Proteins in the sample were digested with L-(tosylamido-2-phenyl) ethyl chloromethyl ketone (TPCK)-treated trypsin (a 1:50 enzyme:substrate ratio; Promega, Madison, WI, USA) at 37 °C for 16 h. To stop the reaction, 25 µl of 0.1% formic acid was added, and the sample was simultaneously spiked with 600 fmol of tryptic digests of Escherichia coli β -galactosidase standard (AB SCIEX, Framingham, MA, USA). The digests were desalted with a C-18 spin column (Thermo), dried via vacuum centrifugation and stored at $-25\,^{\circ}$ C until use.

Selection of ALDH isozyme peptide targets. We gathered the aminoacid sequences of 19 ALDH isozymes from the Uniprot database (released as of 2014.02), digested the sequences in silico with trypsin and selected unique peptides for each ALDH isozyme. Peptides consisted of 6~20 amino acids and had no missing cleavage sites. Detailed selection criteria are described in our previous paper. We considered at least five multiple reaction-monitoring (MRM) transitions of each peptide from spectral libraries such as GPMDB, PeptideAtlas and NIST. If a peptide was available, we used a fragmentation spectrum prediction program, PeptideArt. All candidate peptides were prescreened from six lung cell lines, namely, A549, H23, H322M, H460, IMB90 and Primary, via LC-MRM under the chosen MRM transition parameters. Among these candidates, 28 peptides representing 16 ALDH isozymes were detected in this preliminary test (Table 2). If more than one peptide was detected for a single isozyme, the peptide showing the highest signal was chosen as a representative peptide and was used for quantitation of the corresponding ALDH isozyme in the study.

Liquid chromatography MRM mass spectrometry. Dried tryptic peptides were reconstituted with 30 µl of 5% acetonitrile/0.1% formic acid, injected with a full-sample loop injection of 1 µl and separated in a Nano cHiPLC ReproSil-Pur C18 column (75 μm inside diameter (i.d.) \times 15 cm length, pore size 120 Å, particle size 3 μ m; #804-00011, Eksigent Technologies, Dublin, CA, USA) pre-equilibrated with 95% Solvent A (0.1% formic acid in water) and 5% Solvent B (0.1% formic acid in acetonitrile). Peptides were eluted at a flow rate of 300 nl min⁻¹, with a gradient of 5–10% Solvent B for 4 min, 10-25% for 30 min and 25-60% for 3 min, followed by a 3-min isocratic elution with 60% Solvent B. MRM measurements were performed using a QTrap5500 equipped with a nanoelectrospray ion source (AB SCIEX). The mass spectrometry was operated in the positive mode with the following settings: ion spray voltage of 2100 V, curtain gas at 20 p.s.i., nebulizer gas at 25 p.s.i., resolution at 0.7 Da (unit resolution) for Q1/Q3, interface temperature at 150 °C and a scan mass range of $300 - 1250 \, m/z$. The declustering potential (DP) and collision energy (CE) values were set based on the Skyline (version 2.5) default value. A scheduled MRM mode with detection windows of 300 s and a cycle time of 1.5 s was used. Each sample was analyzed three times with three technical replicates.

Quantification of MRM measurements. Skyline was used to quantify MRM measurements by calculating the peak areas of extracted ion chromatograms. To determine whether a peptide was detected in each cell line, we applied highly stringent criteria: (i) raw peak area greater than 5000, (ii) signal to noise ratio higher than 3 and (iii) coefficient of variation lower than 25% for the technical triplicates. The raw peak area values were normalized as described below.

Two types of standards were used for normalization. Two peptides from human glyceraldehyde-3-phosphate dehydrogenase (Swiss-Prot accession: P04406), GALQNIIPASTGAAK and LISWYDNEFGYSNR, were used as endogenous internal standards and two peptides of *E. coli* β -galactosidase, IDPNAWVER and GDFQFNISR, were used as external spiked standards. Normalized peak areas of target ALDH isoforms were calculated as follows:

$$\check{P}_{i,s} = P_{i,s} \times \sqrt{\sqrt{\prod_{j=1}^{2} \frac{\hat{n}_{j}}{n_{j,s}}}} \times \sqrt{\prod_{k=1}^{2} \frac{\hat{x}_{k}}{x_{k,s}}}$$

where $P_{i,s}$ and $P_{i,s}$ are the normalized and raw peak areas, respectively, of the i-th peptide in sample s, \hat{n}_j is the maximum peak area of the j-th endogenous standard peptide over all samples in the study, $n_{j,s}$ is the raw peak area of the j-th endogenous standard peptide in sample s, \hat{x}_k is the maximum peak area of the k-th exogenous spiked standard peptide over all samples and $x_{k,s}$ is the raw peak area of the k-th exogenous spiked standard peptide in sample s (Table 3). All statistical data were analyzed using the R software (version 2.8.1) and Excel 2010 (version 14.0, Microsoft Office, Redmond, WA, USA).

Measurement of ATP and NADH/NAD+ levels

Total ATP levels were monitored using a CellTiter-Glo Luminescent Cell Viability Assay as per the manufacturer's instructions (G7572, Promega, Durham, NC, USA). CellTiter-Glo was added to 1×10^6 cells, which were then placed on an orbital shaker to induce cell lysis. The samples were then read on a chemiluminescence plate reader (VICTOR light 1420; integration time of 1 s). Total NADH and NAD levels were measured using the NAD/NADH Quantitation Colorimetric Kit (K337-100, BioVision, Milpitas, CA, USA). To obtain total NAD (NADH and NAD), NADH/NAD Extraction Buffer was added to 2×10^5 cells, and a portion of the NADH/NAD extract was heated for 30 min at 60 °C to decompose the NAD. The NADH and NAD extracts were converted to NADH using the NADH-cycling enzyme, and samples were then read on an ELISA reader (Power wave HT, Biotek, Winooski, VT, USA).

FITC Annexin V apoptosis detection

NSCLC cells were treated as indicated for 48 h, washed twice in cold PBS, centrifuged at 1400 r.p.m. for 3 min and resuspended in binding buffer at a concentration of 1×10^6 cells per ml. Portions (100 µl) of the solutions were then transferred to 5-ml culture tubes (1×10^5 cells) before the addition of 5 µl each of annexin V-fluorescein isothiocyanate (FITC) and propidium iodide. Cells were gently vortexed and incubated for 15 min at room temperature (RT) in the dark. Binding buffer (400 µl) was added to each sample before analysis via flow cytometry (BD Biosciences, San Jose, CA, USA).

DNA FACS analysis

A549 and EKVX cells were trypsinized and fixed in 70% (vol/vol) ethanol for DNA staining. For the cell cycle analysis, fixed cells were washed with PBS, and subsequently resuspended in a propidium iodide /RNase staining solution (0.05 mg ml⁻¹ propidium iodide and

0.1 mg ml⁻¹ RNase A in PBS). Fluorescence-activated cell sorting (FACS) analyses were performed by using a FACSCanto II system (BD Biosciences).

Mitotic index

Condensed chromosomes in mitotic cells were intensively visualized by staining them with an aceto-orcein solution (Merck, Billerica, MA, USA) under a light microscope. Mitotic indices were determined based on the percentage of mitotic cells with condensed chromosomes.

Cell cycle synchronization and drug treatment

For the analysis of the G1/S transition, A549 and EKVX cells were blocked at the G1/S boundary by treating them with 2 mm thymidine for 12 or 22 h, respectively, and the cells were treated with gossypol and/or phenformin 6 h before thymidine release. Then, synchronized cells were released into the S phase by washing them three times with PBS. Released cells were incubated again with fresh media plus the same drug(s) they were previously treated with. For the analysis of the G2/M transition, A549 and EKVX cells were blocked at the G1/S boundary by treating them with 2 mm thymidine for 12 or 22 h, respectively, and then releasing them into the S phase by washing the cells three times with PBS and subsequently incubating them with fresh media plus gossypol and/or phenformin.

Flow cytometry

Cells were collected into single-cell suspensions using trypsin/EDTA, fixed with 4% formaldehyde and then permeabilized with 0.1% Triton X-100. Cells were stained with an anti-ALDH1A3 polyclonal antibody and a phycoerythrin-conjugated anti-rabbit antibody (Biotium, Hayward, CA, USA). Flow cytometric analyses were performed at the Flow Cytometry Core (National Cancer Center) using a FACSVerse flow cytometer (BD Biosciences).

Aldefluor activity assay

To measure Aldefluor activity, we used an ALDEFLUOR kit (StemCell Technologies, Vancouver, BC, Canada) according to the manufacturer's instructions. Briefly, cells were suspended in Aldefluor assay buffer containing an ALDH substrate (BODIPY-aminoacetaldehydediethyl acetate, BAAA-DA) and incubated at 37 °C for 45 min. As a negative control for each sample of cells, an aliquot of the sample was treated with diethylaminobenzaldehyde, a specific ALDH inhibitor. The cells were detected in the FL-1 channel of a flow cytometer (BD Biosciences).

Immunohistochemical staining and evaluation

Tissue arrays (CC5, various human lung cancer tissues; CCN5, normal human lung tissues) were purchased from SuperBioChip (Seoul, Korea). Immunohistochemical staining was performed using an UltravisionLP Detection System (Thermo Fisher Scientific Inc., Fremont, CA, USA) and the results are shown in Figure 1. Briefly, after deparaffinization of formalin-fixed, paraffin-embedded human NSCLC tissues, antigens were retrieved in 10 mm citrate buffer (pH 6.0) containing 0.05% Tween-20. The tissues were sequentially treated with 3% hydrogen peroxide and Ultra V blocking solution. After a 1-h incubation at RT with primary antibodies, the slides were washed in Tris-buffered saline with Tween-20 (TBST), incubated with a primary antibody enhancer for 10 min and exposed to horseradish peroxidase-conjugated secondary antibodies for 15 min. After re-washing in TBST, the tissue slides were incubated with the chromogen

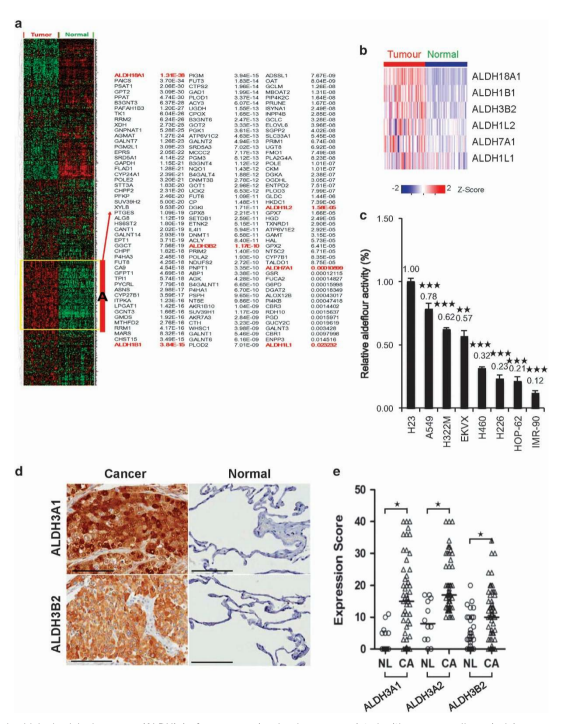


Figure 1 High aldehyde dehydrogenase (ALDH) isoform expression levels are associated with poor overall survival in non-small-cell lung cancer (NSCLC). (a) Metabolic gene clustering of samples from lung adenocarcinoma patients from The Cancer Genome Atlas (TCGA). The clustered genes in set A were upregulated in the tumor group, and 150 metabolic genes with P < 0.05 are listed. (b) Clustering analysis of six ALDH isoforms. Data for ALDH18A1 ($P = 1.31 \times 10^{-38}$), ALDH1B1 ($P = 3.84 \times 10^{-15}$), ALDH3B2 ($P = 1.17 \times 10^{-10}$), ALDH1L2 ($P = 1.58 \times 10^{-05}$), ALDH7A1 ($P = 1.09 \times 10^{-4}$) and ALDH1L1 ($P = 2.32 \times 10^{-2}$) are shown. Gene expression values were normalized using standard normalization (mean = 0 and s.d. = 1). Each row of the heat map is sorted in an ascending order according to the P-value. (c) Relative Aldefluor activity from various NSCLC cell lines and primary human lung cells. **P < 0.01, ***P < 0.001. (d) Immunohistochemical staining of ALDH isoforms in normal and cancerous lung tissue. Scale bar = 100 μm. (e) Expression of ALDH3A1, ALDH3A2 and ALDH3B2 in cancerous (CA, triangles) and normal lung type I and II pneumocytes (NL, circles). *P < 0.001, n = 59 for each sample.

diaminobenzidine (Scytek Laboratories Inc., Logan, UT, USA) and counterstained with Mayer's hematoxylin (Dako Cytomation, Glostrup, Denmark). To evaluate cytoplasmic staining for ALDH isoforms, staining intensity was scored on a scale of 0–3 as follows: 0, no staining of cancer cells; 1, weak staining; 2, moderate staining; 3, strong staining. In addition, the percentage of positive cells among the cancer cells was scored. These two scores were multiplied, and the resulting value was used as the expression score.

Statistical analysis

Statistical analysis was performed using Student's *t*-tests as appropriate.

RESULTS

ALDH isoforms are highly expressed in LUAD

A bioinformatics analysis was performed to assess the expression profiles of metabolic enzymes in NSCLC patients and to identify a distinguishable metabolic context upon which cancer cells depend for survival. The heat map generated from this analysis consists of ~1,600 metabolic genes in 57 pairs of sequencing data selected from 488 LUAD patients (Figure 1a). Our robust clustering analysis and test of statistical significance from the RNA sequencing data for LUADs from The Cancer Genome Atlas revealed several ALDH isoforms that were upregulated in LUAD patients (Figure 1b). The clustering of samples from LUAD patients consistently segregated them into tumor and normal groups (Table 1). ALDH activity in the NSCLC cell lines was measured using an Aldefluor activity kit and was found to be increased in most cell lines compared with activity in control IMR-90 fibroblasts (Figure 1c). To test whether increased expression of ALDH isoforms is also observed in NSCLC cases, immunohistochemical staining was performed for two ALDH subtypes, ALDH3A1 and ALDH3B2.

Table 1 Summary of *P*-values from expression profiling of ALDH isoforms in LUAD patients

Gene name	LUAD (11)
ALDH1A1	1.079E-13
ALDH1A2	3.467E-13
ALDH1A3	0.62022
ALDH1B1	3.84E-15
ALDH1L1	0.023232
ALDH1L2	1.58E-05
ALDH2	6.72E – 17
ALDH3A1	0.18286
ALDH3A2	7.952E-05
ALDH3B1	2.201E-10
ALDH3B2	1.17E-10
ALDH4A1	0.59241
ALDH5A1	0.16429
ALDH6A1	0.7336
ALDH7A1	0.000109
ALDH9A1	0.17913
ALDH18A1	1.31E-38

Abbreviations: ALDH, aldehyde dehydrogenase; LUAD, lung adenocarcinoma.

Other cellular components of normal lung tissue, including type I and II pneumocytes, showed little to no ALDH expression. Lung cancer tissues, however, displayed much higher ALDH levels (Figures 1d and e). Taken together, these data indicate that levels of ALDH isoforms are increased in lung cancer.

To examine their differential expression in NSCLC cells, the expression levels of 16 ALDH isoforms in nine lung cancer cell lines were measured via liquid chromatography MRM mass spectrometry (Figure 2a). To compare the expression levels of each ALDH isoform in the different cell lines, MRM peak areas were normalized against the largest nine values (Figure 2b, Tables 2 and 3). Immunoblotting confirmed the differential expression of these ALDH isoforms (Figure 2c). Flow cytometric analysis of ALDH1A3 showed that this protein is expressed ubiquitously rather than only in a specific cell population, suggesting that ALDH isoforms are widely expressed in NSCLC cells (Figure 2d).

Treatment with gossypol significantly decreases ATP production in NSCLC cells

A total of 15 ALDH isozymes were cloned and expressed in EKVX cells. Our results demonstrate that the transfection of ALDH isozymes generally increases ATP levels by over 10% (Figure 3a). Transfection of ALDH1L1, ALDH1L2, ALDH2, ALDH3A1, ALDH3A2, ALDH3B1 and ALDH18A1 resulted in increases of ATP of over 20%. In particular, ALDH1L1 gene transfection was associated with a 30% increase in ATP production (Figure 3a). Treating each ALDH gene transfectant with 10 µM gossypol, a pan-ALDH inhibitor, reduced ATP production to ~60-70% that of the control, suggesting that gossypol can inhibit most ALDH isoforms in NSCLC cells. Taken together, these data suggest that ALDH has a crucial role in NADH production in the cytoplasm, which is required for ATP production. The effect of gossypol treatment on ATP production in NSCLC cells was compared with its effect on primary lung epithelial cells (Figure 3b). Treating NSCLC cell lines with 10-µM gossypol inhibited ATP production by 25-50% (Figure 3b). Although normal primary lung epithelial cells showed decreases in ATP production of ~28%, the ATP levels in normal cells were four to sevenfold lower than the levels in cancer cells (Figure 3b).

Gossypol induces cell cycle arrest through ATP depletion in NSCLC

Some reports have shown that decreases in ATP levels can induce cell cycle arrest at the G1/S transition as well as at the G2/M transition.^{6,7} To test whether a dual treatment with gossypol and phenformin could be used to regulate the cell cycle, we performed a cell cycle analysis with or without drug treatments. An analysis of cell cycle progression using thymidine block-induced synchronized cells clearly revealed cell cycle progression delay at the G2/M transition (Figure 4a), which agrees with previous reports on the presence of ATP-dependent cell cycle checkpoints at the G2/M transition.^{6,7} Moreover, a delay in mitotic progression was even more

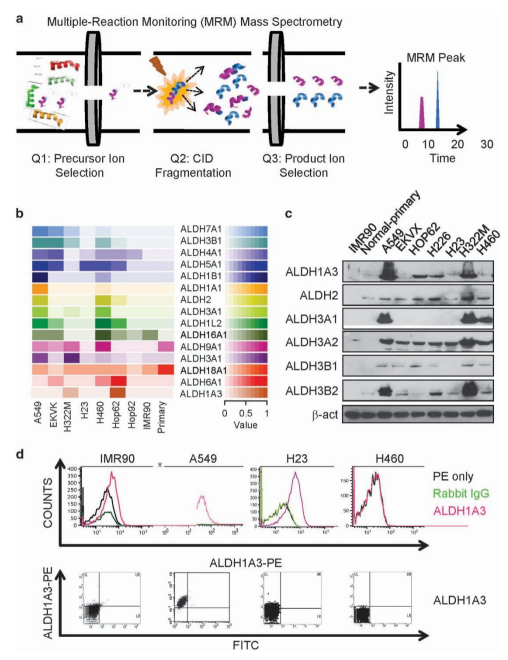


Figure 2 Multiple isoforms of aldehyde dehydrogenase (ALDH) show increased expression in non-small-cell lung cancer (NSCLC) cells. (a) Schematic diagram of liquid chromatography multiple reaction-monitoring mass spectrometry (LC-MRM-MS). (b) Expression of ALDH isoforms in lung cancer cell lines was measured via MRM-MS. (c) Immunoblot analysis of ALDH isoforms in NSCLC cell lines. IMR-90 and normal lung airway cells were used as non-cancer controls, whereas β-actin was used as a loading control. (d) ALDH1A3 was ubiquitously expressed in NSCLC cells as determined via flow cytometric analysis.

significant in the case of the combined treatment (Figure 4a). The inhibition of the G2/M transition seemed to induce significant decreases in mitotic indices in asynchronous cultures of both cell lines (Figure 4b). Further analysis of the cycle of synchronized cells revealed a delay in the cell cycle progression at the G1/S transition in A549 cells as well (Figure 4c), agreeing with previous reports on the presence of ATP-dependent cell cycle checkpoints at the G1/S transition.^{6,7} To test whether the observed cell cycle arrest is due to cell death, cell death was analyzed in A549 and EKVX

cells via FACS at the point of cell cycle arrest (Figure 4d). However, cell death was not detected at the time of cell cycle arrest. This suggests that prolonged cell cycle arrest may stress the cancer cells, eventually inducing apoptosis.

The combination treatment of gossypol and phenformin for 48 h increased cell death by 4.75-fold in EKVX cells compared with cell death in the control treatment, whereas treatment with only gossypol increased cell death by 2.58-fold in EKVX cells compared with cell death in the control treatment (Figure 5a). Moreover, ATP production was decreased by

Table 2 Tryptic peptides of ALDH isozymes detected via MRM-MS

Gene name	Target or standard	Uniprot ID	Species	Retention time (min)	Q1 m/z peptide	CE (volt)	Q3 ions	Top Q3 ion
ALDH1A1	Target	P00352	Homo sapiens (human)	37.8	857.9271 LYSNAYLNDLAGCIK	39.7	y4,y5,y8,y9,y10	9γ
ALDH1A1	Target	P00352	H. sapiens (human)	38.1	772.8909 TIPIDGNFFTYTR	36.7	y5,y8,y9,y10,y11 (2+)	y11(2+)
ALDH1A1	Target	P00352	H. sapiens (human)	36.1	795.4016 ANNTFYGLSAGVFTK	37.5	y3,y7,y9,y10,y11	y10
ALDH1A3	Target	P47895	H. sapiens (human)	33.4	807.8986 ELGEYALAEYTEVK	37.9	y5,y6,y7,y8,y10	у7
ALDH1B1	Target	P30837	H. sapiens (human)	23.8	567.5983 ELGEYGFHEYTEVK	28.5	y8 (2+),y9,y9 (2+),y10,y12	y10(2+)
ALDH1L1	Target	075891	H. sapiens (human)	24.7	714.8483 VLEVEDSTDFFK	34.6	y3,y7,y8,y9,y10	у7
ALDH1L2	Target	Q3SY69	H. sapiens	30.1	786.8863 ANSTEYGLASGVFTR	37.2	y5,y6,y7,y9,y10	99
ALDH2	Target	P05091	H. sapiens (human)	32.7	764.3938 ANNSTYGLAAAVFTK	36.4	y3,y6,y7,y9,y10	99
ALDH2	Target	P05091	H. sapiens (human)	38.7	766.3751 TIPIDGDFFSYTR	36.4	y6,y8,y9,y10,y11 (2+)	y11(2+)
ALDH2	Target	P05091	H. sapiens (human)	30	800.3987 ELGEYGLQAYTEVK	37.7	y4,y6,y7,y9,y10	99
ALDH3A1	Target	P30838	H. sapiens (human)	18.1	549.817 IQQLEALQR	28.7	y2,y4,y5,y6,y7	у7
ALDH3A1	Target	P30838	H. sapiens (human)	20.4	747.883 FDHILYTGSTGVGK	35.8	y8,y9,y10,y11,y12 (2+)	y10
ALDH3A1	Target	P30838	H. sapiens (human)	37.2	724.3807 SLEEAIQFINQR	34.9	y3,y4,y5,y6,y7	у6
ALDH3A2	Target	P51648	H. sapiens (human)	27	500.8055 ILSLLEGQK	26.9	y3,y4,y5,y6,y7	у7
ALDH3A2	Target	P51648	H. sapiens (human)	16.4	633.8017 IAFGGETDEATR	31.7	y6,y8,y9,y10,y10 (2+)	99
ALDH3A2	Target	P51648	H. sapiens (human)	31	716.3902 YIAPTVLTDVDPK	34.6	y10,y10 (2+),y11,y11 (2+)	y10(2+)
ALDH3B1 ^a	Target	P43353	H. sapiens (human)	17.6	776.855 FYGDDPQSSPNLGR	36.8	y5,y6,y7,y9,y10	99
ALDH4A1	Target	P30038	H. sapiens (human)	24.9	565.8245 SSPSLTILAGGK	29.2	y5,y7,y8,y10,y10 (2+)	y5
ALDH5A1	Target	P51649	H. sapiens (human)	32.3	492.8137 LAGLSAALLR	26.6	y3,y4,y5,y6,y8	у6
ALDH5A1	Target	P51649	H. sapiens (human)	23	945.4657 VGNGFEEGTTQGPLINEK	42.9	y3,y6,y7,y9,y11	у3
ALDH6A1 ^a	Target	Q02252	H. sapiens (human)	24.2	597.2935 TLADAEGDVFR	30.4	y3,y5,y7,y8,y9	99
ALDH7A1	Target	P49419	H. sapiens (human)	24.6	952.9622 GEVITTYCPANNEPIAR	43.2	y4,y9,y10 (2+),y14 (2+),y16 (2+) y9	
ALDH9A1	Target	P49189	H. sapiens (human)	13.1	543.3012 EVNLAVQNAK	28.4	y4,y5,y6,y7,y8	у6
ALDH9A1	Target	P49189	H. sapiens (human)	23.2	564.7985 IGDPLLEDTR	29.2	y4,y5,y7,y7 (2+),y8	y7(2+)
ALDH9A1	Target	P49189	H. sapiens (human)	24.8	622.3321 VTIEYYSQLK	31.3	y4,y5,y6,y7,y8	у6
ALDH16A1	Target	Q81Z83	H. sapiens (human)	17.8	570.8223 EALLVANGTPR	29.4	y4,y5,y6,y7,y8	у6
ALDH18A1	Target	P54886	H. sapiens (human)	35	592.8479 GPVGLEGLLTTK	30.2	y3,y4,y6,y7,y9	99
ALDH18A1	Target	P54886	H. sapiens (human)	26.1	478.7981 LNSLAIGLR	26.1	y3,y4,y5,y6,y7	y5
B-gal	External spiked	P16278	Escherichia coli	22.9	550.2802 IDPNAWVER	23.31	y4,y5,y6,y7,y7 (2+)	y7 (2+)
	Standard							
B-gal	External spiked	P16278	E. coli	26.2	542.2645 GDFQFNISR	26.2	y4, y5	у5
	Standard	707	1	L		7	() () () () () () () () () ()	ç
G3F	Endogenous Internal tandard	P04406	<i>H. saplens</i> (numan)	26.5	706.3989 GALGINIIPAS IGAAR	26.41	y4,y5,y6,y8,y9	yx
0	ווופרוומו ומווטמנט			0		r C		Ļ
G3P	Endogenous	P04406	<i>H. sapiens</i> (human)	36.2	882.4049 LISWYDNEFGYSNR	35.91	y3,y5,y6,y9,y12 (2+)	y5
	Internal standard							
Abbroitoito de A		L 0 00000	ON MON	, z z z z z z z z z z z z z z z z z z z	, s + 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0			

Abbreviations: ALDH, aldehyde dehydrogenase; CE, collision energy; MRM-MS, multiple reaction-monitoring mass spectrometry.
^aAnnotated as mitochondrial proteins.

Table 3 Normalized peak areas for the tryptic peptides of 16 ALDH isozymes

	-	;		•								
ALDH		Fragment	Representative									
isozymes	Peptide sequence (Q1)	ion (Q3)	peptide	A549	EKVK	H322M	H23	H460	Hop62	Hop92	IMB90	Primary
ALDH1A1	ANNTFYGLSAGVFTK	y10	z	2 299 577.2	115 108.5	N.D.	N.D.	389 508.8	N.D.	N.D.	N.D.	N.D.
ALDH1A1	LYSNAYLNDLAGCIK	99	z	185 743.7	N.D.ª	N.D.	N.D.	N.D.	N.D.	N.D.	N.D.	N.D.
ALDH1A1	TIPIDGNFFTYTR	y11 (2+)	>-	19 672 834.4	1 106 049.4	N.D.	N.D.	3 193 847.9	N.D.	N.D.	N.D.	12 808.3
ALDH1A3		у7	>-	N.D.	N.D.	54 923.5	N.D.	N.D.	160 067.6	N.D.	N.D.	N.D.
ALDH1B1	ELGEYGFHEYTEVK	y10 (2+)	>-	3 641 491.2	171 602.5	N.D.	N.D.	744 216.5	N.D.	N.D.	N.D.	N.D.
ALDH1L1	VLEVEDSTDFFK	у7	>-	210 455.5	182 277.4	72 664.4	45 930.7	28 234.8	N.D.	N.D.	N.D.	N.D.
ALDH1L2	ANSTEYGLASGVFTR	99	>-	64 268.2	26 451.2	N.D.	N.D.	83 066.2	46 502.7	N.D.	N.D.	N.D.
ALDH2	ANNSTYGLAAAVFTK	99	z	187 727.6	N.D.	N.D.	N.D.	197 033.4	50 184.2	N.D.	N.D.	81 064.3
ALDH2	ELGEYGLQAYTEVK	99	z	150 243.8	N.D.	N.D.	N.D.	166 698.5	41 966.2	46 113.5	21 020.0	63 511.1
ALDH2	TIPIDGDFFSYTR	y11 (2+)	>-	870 545.0	N.D.	N.D.	27 259.6	908 994.7	241 858.7	81 402.2	58 593.9	N.D.
ALDH3A1	FDHILYTGSTGVGK	y10	z	262 594.5	N.D.	62883.0	N.D.	418 238.9	N.D.	N.D.	N.D.	7230.6
ALDH3A1	IQQLEALQR	у7	>-	3 587 171.9	N.D.	862 497.1	N.D.	458 8628.0	N.D.	N.D.	N.D.	N.D.
ALDH3A1	SLEEAIQFINQR	ye	z	1377878.2	N.D.	323 073.4	N.D.	2 035 663.8	N.D.	N.D.	N.D.	N.D.
ALDH3A2	IAFGGETDEATR	99	z	561 081.9	57 309.1	1164782.8	79 575.9	207 168.3	144 229.1	N.D.	N.D.	54 731.4
ALDH3A2	ILSLLEGQK	у7	>-	942 509.0	186 310.6	120 1969.0	209 089.2	335 479.5	168 177.9	45 032.4	132 199.9	180 608.7
ALDH3A2	YIAPTVLTDVDPK	y10 (2+)	z	386 319.7	39 480.3	714 527.4	69 454.7	126 169.9	86 059.2	298 737.1	N.D.	121 247.8
$ALDH3B1^\mathtt{b}$	FYGDDPQSSPNLGR	99	>-	62 262.9	61 949.6	27 169.8	N.D.	35 658.5	8255.1	N.D.	N.D.	N.D.
ALDH4A1	SSPSLTILAGGK	y5	>-	231 740.7	275 607.1	N.D.	77 972.9	149 606.3	142 462.3	113616.3	N.D.	N.D.
ALDH5A1	LAGLSAALLR	ye	>-	34067.6	20 330.2	N.D.	19 581.7	20 374.6	13 829.2	N.D.	N.D.	N.D.
ALDH5A1	VGNGFEEGTTQGPLINE K	у3	z	23639.0	16 276.1	32 714.6	N.D.	N.D.	14 859.9	N.D.	N.D.	N.D.
$ALDH6A1^\mathtt{b}$	TLADAEGDVFR	99	>-	N.D.	19 405.4	N.D.	N.D.	24 117.1	69 238.2	N.D.	N.D.	N.D.
ALDH7A1	GEVITTYCPANNEPIAR	99	>-	845 067.1	641 975.1	305 186.4	N.D.	99 549.5	N.D.	16 614.2	N.D.	N.D.
ALDH9A1	EVNLAVQNAK	ye	z	N.D.	N.D.	N.D.	N.D.	92 854.2	N.D.	N.D.	N.D.	N.D.
ALDH9A1	IGDPLLEDTR	y7 (2+)	>-	113 934.0	78 063.6	135 445.2	N.D.	243 143.6	N.D.	N.D.	N.D.	137 450.8
ALDH9A1	VTIEYYSQLK	ye	z	58 798.9	N.D.	59821.9	N.D.	154 409.4	60 882.6	N.D.	N.D.	50 156.9
ALDH16A1	EALLVANGTPR	ye	>-	83 096.3	83 436.6	N.D.	N.D.	164 803.8	665 40.7	48 868.0	954 69.8	N.D.
ALDH18A1	GPVGLEGLLTTK	99	>-	204 825.7	78224.8	200 087.1	205 250.1	177 663.5	20 7802.9	10 6649.8	200 954.6	575 240.1
ALDH18A1	LNSLAIGLR	y5	z	152 660.6	54 133.0	91 117.2	105 595.7	106 207.5	N.D.	77 145.7	123819.1	190 850.7
Abbragintings.		Z	N D not dotominately	9								

Abbreviations: ALDH, aldehyde dehydrogenase; N, no; N.D., not determined; Y, yes. and raw peak area > 5000. a.N.D.under the criteria of signal-to-noise (SN) ratio > 3, coefficient of variation (CV) < 25% and raw peak area > 5000. bAnnotated as mitochondrial proteins.

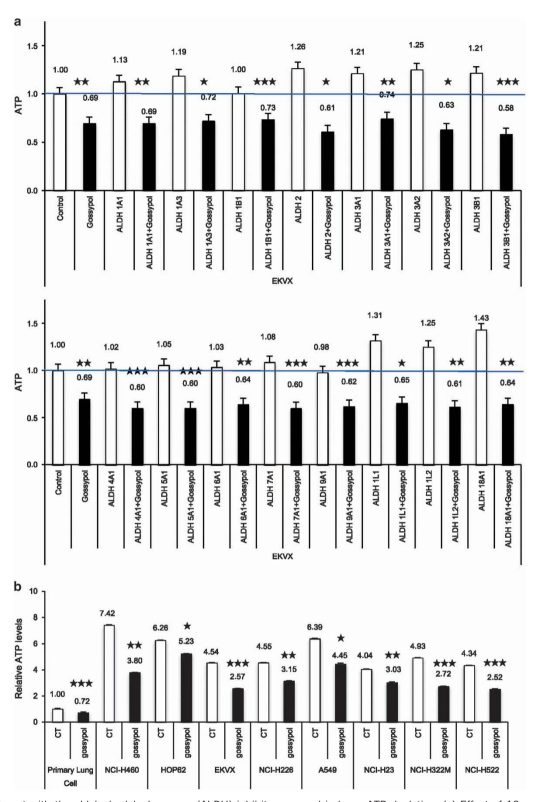


Figure 3 Treatment with the aldehyde dehydrogenase (ALDH) inhibitor gossypol induces ATP depletion. (a) Effect of $10 \, \mu \text{M}$ of gossypol on ATP production in EKVX cells overexpressing various ALDH isoforms. Overexpression primer sets of ALDH isoforms are summarized in the Table 4. (b) Differences in the effect of $10 \, \mu \text{M}$ of gossypol on ATP production between non-small-cell lung cancer (NSCLC) cell lines and primary lung epithelial cells were tested. *P<0.05, **P<0.01, ***P<0.001 compared with control.

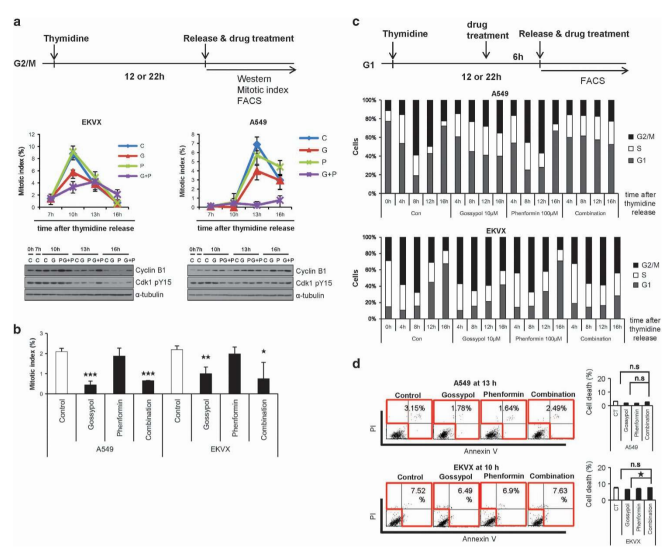


Figure 4 Gossypol combined with phenformin enhances cell cycle arrest and induces cell death in non-small-cell lung cancer (NSCLC). (a) G2/M transition of synchronized cells (following the protocol of a single thymidine block as shown in the upper panel) was monitored via mitotic index assessment (middle panel) and western blot analysis (lower panel) in the presence or absence of the indicated chemicals. Cyclin B1 is a marker for mitotic exit. pY15 of Cdk1 is a marker for the G2/M transition. (b) Effect of gossypol (10 μm), phenformin (100 μm) or a combined treatment for 24 h on the mitotic index of asynchronous cells. *P<0.05, *P<0.01, **P<0.001 compared with vehicle control. (c) G1/S transition of synchronized cells (following the protocol of a single thymidine block as shown in the upper panel) was monitored via DNA FACS analysis (lower panel) in the presence or absence of the indicated chemicals. Note that EKVX cells were mainly arrested at the early S phase, whereas A549 cells were mainly arrested at the G1 phase. (d) To measure cell death, A549 and EKVX cells were treated with gossypol (10 μm), phenformin (100 μm), combination for 13 and 10 h, respectively. Cell death was measured by flow cytometric analysis. Data are representative of the mean and s.d. of three independent experiments. *P<0.05 compared with control.

81% in EKVX cells. We also tested whether gossypol reduced the levels of B-cell lymphocyte/leukemia-2 (Bcl-2) during the induction of cell death and found that the treatment resulted in no change in Bcl-2 levels (Figure 5b).

DISCUSSION

This study revealed that gossypol combined with phenformin significantly increases the therapeutic response of NSCLC. Here, we found that gossypol induces cell cycle arrest via ATP depletion, which can produce greater anticancer effects when combined with phenformin treatment.

ALDH expression has been recognized as a drug resistance marker in cancer (see reviews in Januchowski *et al.*⁸). By exploring the role of ALDH, previous studies have found that the expression of ALDH isoforms such as ALDH1 and ALDH3A1 are highly upregulated in cancer stem cells.⁹ A proposed role of ALDH in cancer stem cells is the conversion of activated cyclophosphamide to the inactive excretory product carboxyphosphamide.¹⁰ Here, we found that ALDH isoforms are expressed in every cell of cancer cell lines, which conflicts with cancer stem cell theory. However, a previous report agrees with our observation that ALDH1 is not a suitable

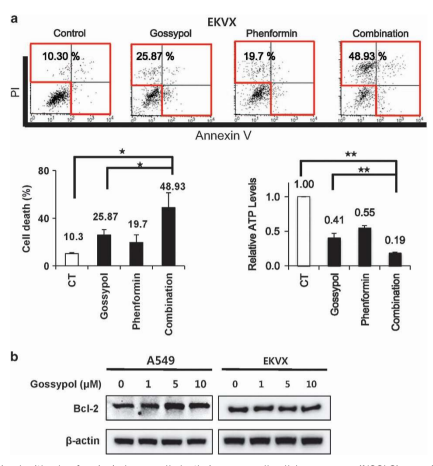


Figure 5 Gossypol combined with phenformin induces cell death in non-small-cell lung cancer (NSCLC) associated with ATP depletion. (a) Effect of gossypol ($10 \,\mu\text{M}$), phenformin ($100 \,\mu\text{M}$) or a combined treatment for 48 h on cell death as determined via flow cytometric analysis and ATP production determined via ATP assay. (b) Effect of gossypol ($10 \,\mu\text{M}$), phenformin ($100 \,\mu\text{M}$) or a combined treatment on Bcl-2 expression in NSCLC cells was determined via immunoblotting. Data are representative of the mean and s.d. of three independent experiments. *P<0.05, **P<0.01 compared with control.

CSC marker because its expression is also detected in epithelial cancer cell lines. ¹¹ Therefore, there is no clear evidence for the role of ALDH in cancer, although ALDH inhibitors increase the effect of chemotherapy. ¹² This study revealed that NSCLC is highly dependent on ALDH for ATP production because ALDH produces NADH, which acts as an electron carrier for ATP synthesis.

Gossypol is a terpenoid extracted from cottonseeds that has been used as an antispermatogenic agent in Chinese medicine. Gossypol is known as a non-competitive inhibitor of substrate binding in ALDH-induced oxidation and a strong competitive inhibitor against cofactors such as NAD+. ^{13,14} *In vitro* kinetic studies have shown that gossypol exerts greater effects on the ALDH3 isozyme than on the ALDH1 and ALDH2 isozymes¹⁵ (see reviews in Kopakka *et al.*¹⁴). Gossypol has also been identified to competitively bind to Bcl-2 homology (BH) domains in Bcl-2 family, which blocks the BH3 dimerization domain of pro-apoptotic proteins binding to Bcl-_{XL}. ¹⁶ In line with this theory of Bcl-2 inhibition, gossypol treatment has been shown to decrease Bcl-2 protein levels. ^{17,18} However, in this study, we did not observe any change of Bcl-2 protein levels (Figure 5b). Whatever the mechanism, single or

combined treatments with gossypol have been associated with limited anticancer efficacy in clinical trials. Treatment with gossypol alone (40 mg per day) during a phase I/II clinical study of metastatic breast cancer refractory to doxorubicin and paclitaxel also demonstrated a limited therapeutic response. Recently, a randomized study combining AT-101 (R-(-)-gossypol acetic acid) with docetaxel in patients with advanced NSCLC showed negligible survival advantages as a second-line treatment. This suggests that there is a limit to the use of gossypol as a single therapeutic option in treating cancer.

The Warburg effect in cancer proposes that glucose is the primary source of carbon for biomolecule synthesis²¹ even though glycolysis contributes to ATP synthesis.

In this study, we found that ALDH contributes to ATP production through the supply of NADH in NSCLC. This suggests that NSCLC depends on cytosolic NADH for ATP production in the mitochondria. However, the mechanisms involved in NADH transportation via the malate–aspartate shuttle remain to be verified in NSCLC.

Phenformin is known to block electron transfer at mitochondrial complex I, which results in the inhibition of ATP synthesis. The combined treatment with gossypol and

Table 4 Expression constructs of ALDH isoforms used in this study

				Insert		
No	Vector	Gene template		Primer sequences	Enzyme site	Amplicon size (bp)
1	p3XFlag-CMV	ALDH1A1	F'	TGACGATGACAAGCTTATGTCATCCTCAGGCACG	<i>Hin</i> dIII	1506
			R′	TCGCGGCCGCAAGCTT TTATGAGTTCTTCTGAGAG	<i>Hin</i> dIII	
2	p3XFlag-CMV	ALDH1A3	F′	TGACGATGACAAGCTT ATGGCCACCGCTAACGGGGC	<i>Hin</i> dIII	1539
			R′	TCGCGGCCGCAAGCTT TCAGGGGTTCTTGTCGCC	<i>Hin</i> dIII	
3	p3XFlag-CMV	ALDH1B1	F′	TGACGATGACAAGCTT ATGCTGCGCTTCCTGGCAC	<i>Hin</i> dIII	1554
			R′	TCGCGGCCGCAAGCTT TTACGAGTTCTTCTGAGG	<i>Hin</i> dIII	
4	p3XFlag-CMV	ALDH2	F′	TGACGATGACAAGCTT ATGTTGCGCGCTGCCGCC	<i>Hin</i> dIII	1554
			R′	TCGCGGCCGCAAGCTT TTATGAGTTCTTCTGAGGC	<i>Hin</i> dIII	
5	p3XFlag-CMV	ALDH3A1	F′	TGACGATGACAAGCTT ATGAGCAAGATCAGCGAG	<i>Hin</i> dIII	1362
			R′	TCGCGGCCGCAAGCTT TCAGTGCTGGGTCATCTTG	<i>Hin</i> dIII	
6	p3XFlag-CMV	ALDH3A2	F′	TGACGATGACAAGCTT ATGGAGCTCGAAGTCCGGC	<i>Hin</i> dIII	1527
			R′	TCGCGGCCGCAAGCTT TCATCTCTGCTTACTGGAC	<i>Hin</i> dIII	
7	p3XFlag-CMV	ALDH3B1	F′	TGACGATGACAAGCTT ATGGACCCCCTTGGGGAC	<i>Hin</i> dIII	1407
			R′	TCGCGGCCGCAAGCTT TCAGAGCAGTGTGCAGCTG	<i>Hin</i> dIII	
8	p3XFlag-CMV	ALDH4A1	F′	TGACGATGACAAGCTT ATGCTGCTGCCGGCGCCCCG	<i>Hin</i> dIII	1692
			R′	TCGCGGCCGCAAGCTT TCACTGCATGTACGCGTAG	<i>Hin</i> dIII	
9	p3XFlag-CMV	ALDH5A1	F'	TGACGATGACAAGCTT ATGGCGACCTGCATTTGGC	<i>Hin</i> dIII	1647
			R′	TCGCGGCCGCAAGCTT CTACAAGCCCCCGTAACAC	<i>Hin</i> dIII	
10	p3XFlag-CMV	ALDH6A1	F'	TGACGATGACAAGCTT ATGGCGGCGCTATTGGCG	<i>Hin</i> dIII	1608
			R′	TCGCGGCCGCAAGCTT CTAACGGCCCATGGTAGGC	<i>Hin</i> dIII	
11	p3XFlag-CMV	ALDH7A1	F'	TGACGATGACAAGCTT ATGTGGCGCCTTCCTCGC	<i>Hin</i> dIII	1620
			R′	TCGCGGCCGCAAGCTT TTACTGAAACTTGATTCC	<i>Hin</i> dIII	
12	p3XFlag-CMV	ALDH9A1	F′	TGACGATGACAAGCTT ATGTTTCTCCGAGCAGGC	<i>Hin</i> dIII	1557
			R′	TCGCGGCCGCAAGCTT TCAAAAAGCAGATTCCACATC	<i>Hin</i> dIII	
13	p3XFlag-CMV	ALDH1L1	F′	TGACGATGACAAGCTT ATGAAGATTGCAGTGATTG	<i>Hin</i> dIII	2709
			R′	TCGCGGCCGCAAGCTT TCAGTATTCGAAGGTCAC	<i>Hin</i> dIII	
14	p3XFlag-CMV	ALDH1L2	F′	TGACGATGACAAGCTT ATGCTGCGGCGGGGCAGC	<i>Hin</i> dIII	2772
	-		R′	TCGCGGCCGCAAGCTT CTAATATTCCAGTGTCAC	<i>Hin</i> dIII	
15	p3XFlag-CMV	ALDH18A1	F′	TGACGATGACAAGCTT ATGTTGAGTCAAGTTTAC	<i>Hin</i> dIII	2382
	-		R′	TCGCGGCCGCAAGCTT TCAGTTGGTGTTTCTCTG	<i>Hin</i> dIII	

Abbreviations: ALDH, aldehyde dehydrogenase.

phenformin potentiates the induction of G1 and G2 arrest, which induces cell death in NSCLC cell lines (Figure 5). Therefore, gossypol increases the therapeutic response of NSCLC to phenformin (Figure 5) and may provide a novel therapeutic approach for treating NSCLC.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

ACKNOWLEDGEMENTS

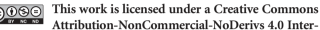
This work was supported by a research grant from the National Cancer Center of Korea to DH, HJ, KMH and SYK (NCC1410670). CL was supported by an NRF Multi-Omics Program (2012M3A9B9036679).

- Wong DW, Leung EL, So KK, Tam IY, Sihoe AD, Cheng LC et al. The EML4-ALK fusion gene is involved in various histologic types of lung cancers from nonsmokers with wild-type EGFR and KRAS. Cancer 2009; 115: 1723–1733.
- 3 Li X, Wan L, Geng J, Wu CL, Bai X. Aldehyde dehydrogenase 1A1 possesses stem-like properties and predicts lung cancer patient outcome. *J Thorac Oncol* 2012; **7**: 1235–1245.
- 4 Jiang F, Qiu Q, Khanna A, Todd NW, Deepak J, Xing L et al. Aldehyde dehydrogenase 1 is a tumor stem cell-associated marker in lung cancer. Mol Cancer Res 2009; 7: 330–338.
- 5 Cancer Genome Atlas Research N, Weinstein JN, Collisson EA, Mills GB, Shaw KR, Ozenberger BA et al. The Cancer Genome Atlas Pan-Cancer analysis project. Nat Genet 2013; 45: 1113–1120.
- 6 Sweet S, Singh G. Accumulation of human promyelocytic leukemic (HL-60) cells at two energetic cell cycle checkpoints. *Cancer Res* 1995; 55: 5164–5167.
- 7 Sweet S, Singh G. Changes in mitochondrial mass, membrane potential, and cellular adenosine triphosphate content during the cell cycle of human leukemic (HL-60) cells. *J Cell Physiol* 1999; 180: 91–96.
- 8 Januchowski R, Wojtowicz K, Zabel M. The role of aldehyde dehydrogenase (ALDH) in cancer drug resistance. Biomed Pharmacother 2013; 67: 669–680.
- 9 Ma I, Allan AL. The role of human aldehyde dehydrogenase in normal and cancer stem cells. Stem Cell Rev 2011; 7: 292–306.

¹ Pao W, Chmielecki J. Rational, biologically based treatment of EGFR-mutant non-small-cell lung cancer. Nat Rev Cancer 2010; 10: 760–774.

- 10 Kastan MB, Schlaffer E, Russo JE, Colvin OM, Civin CI, Hilton J. Direct demonstration of elevated aldehyde dehydrogenase in human hematopoietic progenitor cells. *Blood* 1990; **75**: 1947–1950.
- 11 Deng S, Yang X, Lassus H, Liang S, Kaur S, Ye Q *et al.* Distinct expression levels and patterns of stem cell marker, aldehyde dehydrogenase isoform 1 (ALDH1), in human epithelial cancers. *PLoS ONE* 2010; **5**: e10277.
- 12 Kohn FR, Landkamer GJ, Manthey CL, Ramsay NK, Sladek NE. Effect of aldehyde dehydrogenase inhibitors on the ex vivo sensitivity of human multipotent and committed hematopoietic progenitor cells and malignant blood cells to oxazaphosphorines. *Cancer Res* 1987; **47**: 3180–3185.
- 13 Burgos C, Gerez de Burgos NM, Rovai LE, Blanco A. In vitro inhibition by gossypol of oxidoreductases from human tissues. *Biochem Pharmacol* 1986; **35**: 801–804.
- 14 Koppaka V, Thompson DC, Chen Y, Ellermann M, Nicolaou KC, Juvonen RO et al. Aldehyde dehydrogenase inhibitors: a comprehensive review of the pharmacology, mechanism of action, substrate specificity, and clinical application. *Pharmacol Rev* 2012; **64**: 520–539.
- 15 Rekha GK, Sladek NE. Inhibition of human class 3 aldehyde dehydrogenase, and sensitization of tumor cells that express significant amounts of this enzyme to oxazaphosphorines, by the naturally occurring compound gossypol. *Adv Exp Med Biol* 1997; **414**: 133–146.
- 16 Kitada S, Leone M, Sareth S, Zhai D, Reed JC, Pellecchia M. Discovery, characterization, and structure-activity relationships studies of proapoptotic polyphenols targeting B-cell lymphocyte/leukemia-2 proteins. *J Med Chem* 2003; **46**: 4259–4264.
- 17 Moretti L, Li B, Kim KW, Chen H, Lu B. AT-101, a pan-Bcl-2 inhibitor, leads to radiosensitization of non-small cell lung cancer. *J Thorac Oncol* 2010; **5**: 680–687.

- 18 Ren T, Shan J, Qing Y, Qian C, Li Q, Lu G et al. Sequential treatment with AT-101 enhances cisplatin chemosensitivity in human non-small cell lung cancer cells through inhibition of apurinic/apyrimidinic endonuclease 1-activated IL-6/STAT3 signaling pathway. Drug Des Dev Ther 2014; 8: 2517–2529.
- 19 Van Poznak C, Seidman AD, Reidenberg MM, Moasser MM, Sklarin N, Van Zee K et al. Oral gossypol in the treatment of patients with refractory metastatic breast cancer: a phase I/II clinical trial. Breast Cancer Res Treat 2001; 66: 239–248.
- 20 Ready N, Karaseva NA, Orlov SV, Luft AV, Popovych O, Holmlund JT et al. Double-blind, placebo-controlled, randomized phase 2 study of the proapoptotic agent AT-101 plus docetaxel, in second-line non-small cell lung cancer. J Thorac Oncol 2011; 6: 781–785.
- 21 Guo Q, Liu Z, Jiang L, Hu T, Li D, Liu Y *et al.* Starvation-induced autophagy in cultured non-small cell lung cancer cells. *Nan Fang Yi Ke Da Xue Xue Bao* 2014; **34**: 627–630.



national License. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by-nc-nd/4.0/