Characterization of cancer omics and drug perturbations in panels of lung cancer cells

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To better understand the disruptions of transcriptional regulations and gene expression in lung cancers, we constructed a multi-omics catalogue of the responses of lung cancer cells to a series of chemical compounds. We generated and analyzed 3,240 RNA-seq and 3,393 ATAC-seq libraries obtained from 23 cell lines treated with 95 well-annotated compounds. To demonstrate the power of the created multi-omics resource, we attempted to identify drugs that could induce the designated changes alone or in combination. The basal multi-omics information was first integrated into co-expression modules. Among these modules, we identified a stress response module that may be a promising drug intervention target, as new combinations of compounds that could be used to regulate this module and the consequent phenotypic appearance of cancer cells have been identified. We believe that the multi-omics profiles generated in this study and the strategy used to stratify them will lead to more rational and efficient development of anticancer drugs.

Cancer clinical sequencing analyses at a varying scales identify the genotypes of cancers, including the driver mutations in lung adenocarcinoma cells1-2. These mutations are subjected to further analysis as potential targets for molecular targeting drugs. For example, EGFR tyrosine kinase inhibitors (e.g., gefitinib and erlotinib) and ALK inhibitors (e.g., crizotinib) are administered to patients with EGFR mutations and ALK fusions, respectively3-4. Suitable therapeutic approaches based on the genomic mutation patterns of each cancer have been successful. Nevertheless, approximately 30% of patients with lung adenocarcinoma do not benefit from this approach because appropriate molecular targeting drugs are not available5.

To alleviate shortages in the repertoire of chemical compounds that may serve as “seeds” for eventual drug development, current approaches utilize “high-throughput screening” which is becoming more expensive. Even when starting with millions of compounds, eventual success in obtaining a desirable drug cannot always be guaranteed. To overcome this drawback, the intensive utilization of the accumulated knowledge of genomes, epigenomes, and the transcriptional activities of cells and their molecular networks is currently underway; for example, the Encyclopedia of DNA Elements (ENCODE) database has been used for such studies7.

Indeed, omics information for cell lines, which are frequently used for cancer research, has been rapidly accumulated8-10. By making use of panels of cancer cell lines, numerous studies have revealed the details of the associations among genotypes and phenotypes. For example, the Cancer Cell Line Encyclopedia (CCLE) and other large projects have generated data on the genotypes and cellular responses to pharmacological intervention of hundreds of cancer cell lines10-12. Furthermore, target screening systems were applied to enrich the informational resources for the panels, particularly when novel genetic vulnerabilities in the cancer cells were investigated13-16. These high-throughput screens provide useful information about how the products of individual genes or their combinations could efficiently kill cancer cells and could therefore be targeted by drugs. Moreover, the Connectivity Map (CMAP)17 project compiled a large collection of gene expression profiles of cell lines treated...
Figure 1. High-throughput transcriptome and epigenome perturbations. (a) Reproducibility of the RNA-seq data. The gene expression levels of the A549 cells (DMSO control) were compared. Left panel: the reproducibility among the different plates was evaluated. Right panel: the datasets within the same plates were compared. The Pearson correlation coefficients are shown in the inset. (b) Expression changes in A549 cells treated with CUDC 101. The fold changes in expression were calculated and compared between dataset 1 and dataset 2 for similar treatment conditions (≥4-fold and ≤1/4-fold changes are shown in red and blue, respectively). The Pearson correlation coefficients are shown in the inset. (c) Reproducibility of the ATAC-seq data. The signal intensities reflecting the open chromatin status are shown for the promoter regions of genes in the A549 cells in a manner similar to that shown in (a). (d) Quantification of the changes in the open chromatin status in A549 cells treated with CUDC 101. The fold changes in the intensities of the ATAC-seq signals were compared between the datasets in a manner similar to that shown in (b) (≥2 and ≤1/2-fold changes are shown in red and blue, respectively). (e) Association between the expression levels (RNA-seq) and chromatin accessibility of the promoter regions (ATAC-seq). (f) Comparison of transcriptome profiles between the in-
Evaluation of the RNA-seq and ATAC-seq datasets. We obtained 3,240 data points for the RNA-seq analysis. The RNA-seq data for each compound comprised an average of 1,356,520 and 1,794,226 sequencing reads in dataset 1 and dataset 2, respectively (Table 1). For quality control and initial evaluation purposes, the data were screened for the presence of spike-in controls. The expression values for spike-in controls were validated using qPCR analysis (Fig. 2g). We obtained 3,240 data points for the RNA-seq analysis. The RNA-seq data for each compound comprised an average of 1,356,520 and 1,794,226 sequencing reads in dataset 1 and dataset 2, respectively (Table 1). For quality control and initial evaluation purposes, the data were screened for the presence of spike-in controls. The expression values for spike-in controls were validated using qPCR analysis (Fig. 2g).
mapped reads were required to comprise at least 0.5 million reads containing 15% or fewer intron-mapped reads with an average reads per kilobase million (rpkm) value within ±2 standard deviations (sd) of that of the spike-in control for all wells. After the initial filtration, we further compared the expression abundances (rpkm values) of the protein-coding genes within the datasets to validate the reproducibility of the expression profiles using genes with an rpkm value ≥2 (Figs. 1a and S3). We observed a reasonably high positive correlation between the RNA-seq datasets (r = 0.90, concordance correlation coefficient (ccc) = 0.90 within each plate; r = 0.88, ccc = 0.88 between plates for A549 cells). The data was also reasonably correlated with standard RNA-seq data (Illumina TruSeq) that was previously obtained (r = 0.76 and r = 0.73) (Supplementary Fig. S3c). We also found that the gene expression changes were precisely represented in individual cases. An example of the changes induced by an HDAC inhibitor (CUDC 101) is shown in Fig. 1b. We observed that fold expression changes were moderately correlated between the RNA-seq datasets (r = 0.31, ccc = 0.31). The limited fraction of the genes showed significant changes even under the treatment with epigenetic targeting drugs (see also Supplementary Fig. S4).

For the ATAC-seq analysis, we selected ATAC-seq datasets with >10,000 peaks detected using MACS228, yielding 3,393 ATAC-seq datasets with an average of 8,158,509 sequencing reads. We calculated the tag densities (ppm values) for the promoter and enhancer regions to determine the chromatin accessibility. Similar to the RNA-seq analysis, we evaluated the reproducibility of the ATAC-seq analysis. We found that the tag densities of the promoters were correlated between dataset-1 and 2 under the control condition (r = 0.96, ccc = 0.95 within each plate, r = 0.86, ccc = 0.85 between plates of A549 cells) (Figs. 1c and S3). We examined the changes in open chromatin in drug-treated cells (r = 0.36, ccc = 0.32 for promoters; r = 0.38, ccc = 0.35 for enhancers between the datasets under the CUDC 101 treatment) (Fig. 1d). For the RNA-seq and ATAC-seq data, we also conducted saturation analysis of the sequencing depths to investigate whether the sequencing depths were adequate for high-throughput monitoring of drug perturbation. We confirmed that >1 M reads were enough to analyze the expression patterns in the high-throughput RNA-seq. For the ATAC-seq, the open chromatin patterns and the numbers of the detected promoters and enhancers were saturated at >2 M reads (Supplementary Fig. S5).

When we compared the RNA-seq data and the ATAC-seq data for the promoter regions, we found that they were moderately correlated (Spearman r = 0.36) (Fig. 1e). The inconsistencies between the transcriptome and
epigenome data may indicate that the promoter regions likely included open chromatin regions that lacked enhancer activity under any particular condition. Additionally, other regulatory factors may have affected transcription such as the open chromatin status of the enhancer regions.

For the validation studies, we compared the RNA-seq profiles from this study with the CCLE data (Fig. 1f). To confirm the mode of action, as measured by our procedures, we also evaluated whether the transcriptome signatures represented in our datasets were consistent with the L1000 CMAP signatures. Using the CMAP dataset as a reference, we confirmed that the transcriptome signatures of vorinostat were highly concordant with those of HDAC inhibitors in the CMAP dataset (Fig. 1g). These results suggest that our RNA-seq data accurately represented the transcriptome mode of action for a particular drug. By conducting qPCR validation, we investigated whether the changes in transcriptome and epigenome status during the drug treatments were accurately calculated by the sequencing and post sequencing analyses. The results of the qPCR experiments using the RNA-seq and ATAC-seq libraries were generally highly consistent with the results of the RNA-seq and ATAC-seq analyses (Fig. 1h,i; also see Supplementary Fig. S6). Based on this validation, we considered that the substantial collected datasets could sufficiently represent the cellular changes reflected in the multi-omics profiles in response to the given chemical compounds.

Multi-omics and the phenotypes of the lung adenocarcinoma cell lines. To measure the effects of the drugs on cell survival rates, we used 320 compounds, including those used for the multi-omics perturbation analysis. We measured the viability of each drug-treated cell line (Fig. 2a and Supplementary Table S2). This high-throughput screening assay provided a high average Z' factor of 0.80 (range: 0.59–0.90). Consistent with previous studies, we could evaluate the expected correlations between the genomic mutations/multi-omics statuses and the sensitivity of the cells to a series of drugs, including the EGFR inhibitors (Fig. 2b), multi tyrosine kinase inhibitors (i.e., crizotinib and vandetanib) and epigenetically targeted drugs (i.e., BET inhibitors) (Supplementary Fig. S7).

Based on the analysis of 26 representative cancer-related genes, including oncogenes, tumor suppressor genes, chromatin remodeling factors, and oncogenic fusion genes, we divided the cell lines into two groups according to the cellular IC_{50} values and then compared the multi-omics statuses of the 26 genes in the two groups. This analysis identified 17 combinations associated with cell survival and genome and expression signatures (left panel, Fig. 2c). Genomic KEAP1 mutations were significantly enriched in cells sensitive to the MEK inhibitor PD0325901 (middle and right panels, Fig. 2c). In cancer cells, oncogenic MAPK signaling is aberrantly upregulated and may upregulate NFE2L2 transcription. KEAP1 generally regulates the transcription factor NFE2L2 via a ubiquitin proteasome pathway. When KEAP1 is mutated and NFE2L2 signaling is stabilized, cancer cells may be more addicted to MAPK oncogenic and KEAP1-NFE2L2 oxidation reduction (redox) signals and may become sensitive to MEK inhibitors. The identified KEAP1-centered module was subjected to further detailed analyses, as described in the following section (see below).
Figure 4. Antioxidative stress modules. (a) Sensitivity to auranofin was associated with activity of the "lightcyan (putative redox)" module. The IC_{50} values were compared with the module eigengenes of the module. The Spearman correlation coefficient is shown in the inset. (b) NFE2L2 targets were enriched within the "lightcyan (putative redox)" module members. The proportion of genes with NFE2L2 binding sites in the promoter or enhancer is shown for each module. (c) The association between KEAP1-NFE2L2 mutations and "lightcyan (putative redox)" module activity. The mutational patterns of KEAP1 and NFE2L2 are shown in the upper panel. The module activity is shown in the lower panel. The cell lines are ordered according to the ME for the module. (d) Expression patterns of the "lightcyan (putative redox)" module members in 515 TCGA-LUAD cases. The median fold changes in the expression levels are shown in the heat map. The mutational statuses of the KEAP1 and NFE2L2 genes are shown in the side bar in green and orange, respectively. The color key is shown in the margin. (e) Correlation between the amounts of UDP-glucuronate and "lightcyan (putative redox)" module activity. The Spearman correlation coefficient is shown in the inset. (f) NFE2L2 knockdown downregulated "lightcyan (putative redox)" module members. (g) Kaplan-Meier analysis of TCGA-LUAD cases
based on the activity of the “lightcyan (putative redox)” module. Overall survival time was compared between cases with high module activity (asterisk in d) and other cases. (h) The putative redox module. Representative member genes in the “lightcyan (putative redox)” module are shown along the associated metabolic pathways.

We used RNA-seq and ATAC-seq to identify the modes of action of the drugs (Fig. 1). For more detailed characterization of the drugs, we should consider the complex multi-omics status of each patient, which has not been sufficiently considered; this would strongly affect the transcriptional responses to drugs. Below, we describe transcriptome module-based target selection for the rational administration of drugs to lung cancer patients who harbor diverse genomic, epigenomic, and transcriptomic characteristics. We confirmed that potentially informative influences of drug treatments were unrecognized when assessed according to the cell survival rates.

**Gene co-expression network analysis to identify gene expression modules.** To integrate the omics responses with the cellular responses, which remained latent occasionally, we first evaluated the multi-omics profiles according to the transcriptional regulatory modules. Initially, based on the RNA-seq data for all cell lines included in the datasets of untreated conditions, we employed the weighted gene co-expression network analysis (WGCNA) method\(^\text{31}\). We identified 73 co-expression modules within the transcriptional signatures (Fig. 3a, Supplementary Fig. S8 and Supplementary Table S3). We expected that individual genes with similar expression patterns, which were likely regulated by shared mechanisms, would be modulated. Thus, a bird's eye-view analysis of the responses to drug perturbation would be possible. We selected 51 modules for further analyses after excluding 22 modules that were likely included mainly due to cell line-specific copy number aberrations. WGCNA analysis offers a major advantage because the value of the “module eigengene” (ME) may be used as an indicator to represent the activity of the corresponding module (Fig. 3b). The ME is calculated based on the first principal component of the expression patterns of the gene members included in the module.

**The antioxidative stress response module was associated with KEAP1-NFE2L2 mutations.** We particularly focused on the “lightcyan” module, which may be controlled by a specific repertoire of drugs (Fig. 4a, see below). The “lightcyan” module comprised 158 genes, which were significantly associated with redox-related pathways enriched in “GO:0016616 oxidoreductase activity, acting on the CH-OH group of donors, NAD or NADP as acceptor” (Bonferonni \(p = 0.0013\)) (Supplementary Table S4). The “lightcyan” module included genes that respond to oxidative stressors such as GSR and IDH1\(^\text{32}\) (Supplementary Table S3).

To identify the transcription factor(s) controlling this module, we analyzed the ENCODE database\(^\text{31}\) to determine whether transcription factor bindings were enriched within the proximal regions of the 158 member genes. We found that NFE2L2 may represent an associated transcription factor. The binding of NFE2L2 to the promoter or enhancer regions of 67 genes (44% of the “lightcyan” module members) was indicated by the ChIP-seq data from A549 cells in the ENCODE dataset (Fig. 4b). In lung adenocarcinoma cells, either the activation of NFE2L2 or the loss of its negative regulator KEAP1 frequently occur. We found that cell lines harboring KEAP1 or NFE2L2 mutations showed significantly increased module activity as represented by their module eigengene (\(p = 0.0001\), Wilcoxon rank sum test) (Fig. 4c). For clinical specimens, we also retrieved and analyzed the RNA-seq and whole exome sequencing datasets from the TCGA\(^\text{1}\). We analyzed the 515 clinical samples of lung adenocarcinoma (TCGA-LUAD) and found that the activity of the “lightcyan” module was strongly correlated with the mutational status of KEAP1 and NFE2L2 (Fig. 4d).

We then analyzed the functional relevance of this module. NFE2L2 is a transcription factor that is activated by oxidative stress and controls the expression of genes that encode proteins that mediate downstream redox reactions, thereby providing robust activity against oxidative stress in cancer cells\(^\text{35}\). It was also essential to analyze metabolites for understanding these redox reactions. We measured abundances of 113 metabolites by metabolome profiling (see Methods). When we analyzed the metabolic profiles of the cell lines, we found that seven metabolites were significantly correlated with “lightcyan” module activity (Spearman \(r = 0.84\), \(p < 0.002\)). UDP-glucuronate (Spearman \(r = 0.84\), Fig. 4e) serves as a substrate of UDP-glucose 6-dehydrogenase, which is encoded by UGDH, a member of the “lightcyan” module and a transcriptional target of NFE2L2. Further evidence that supports the conclusion that other metabolites are associated with this NFE2L2-centered module is presented in Supplementary Table S5. For modules which were associated with metabolic profiles, such as the “lightcyan” module, the genes encoding enzymes could be identified as the key genes of the modules. For these cases, we found that metabolites are, indeed, good indicators representing functional and biological relevance of the modules.

Finally, we conducted an NFE2L2 knockdown experiment in the A549 cell line. We directly confirmed the downregulation of several core members of the “lightcyan” module (Fig. 4f). The lines of evidence collectively indicated that the “lightcyan” module was an NFE2L2-centered module, and the activation of the “lightcyan” module was represented by the transcriptome signatures of KEAP1- and NFE2L2-mutant cancers.

To evaluate the clinical effects of controlling this module, we reanalyzed the TCGA datasets. Among a group of cancers (128 cases, 25%) with high levels of “lightcyan” module activity, significantly poor prognosis was observed (\(p = 0.019\)) (Fig. 4g). We further analyzed other cancer types and identified the same transcriptional dysregulation in cancers such as lung squamous cell carcinoma and esophageal carcinoma (198 and 35 cases; 39% and 19%, respectively) (Supplementary Fig. S9), indicating the general importance of this module in cancer. Therefore, effective drug intervention strategies may have broad clinical relevance.

**Drug interventions in the “lightcyan” module using auranofin.** The activity of the “lightcyan” module was significantly correlated with sensitivity to auranofin treatment (\(r = 0.86\), Spearman correlation...
coefficient) (Fig. 4a). Specifically, cell lines with increased expression levels of “lightcyan” module genes were resistant to auranofin. We sought to determine whether auranofin might be able to regulate the “lightcyan” module. Auranofin is used to treat rheumatoid arthritis. Auranofin is an inhibitor of thioredoxin reductase (encoded by TXNRD1/2)32. This drug is used as an approved drug to treat rheumatoid arthritis.

Figure 4h illustrates the association of antioxidative stress signaling with the “lightcyan” module. Auranofin may be associated with this module by regulating certain target genes of NFE2L2. However, in cell lines with increased “lightcyan” activity, other antioxidative stress genes and auranofin targets were also dysregulated, and these genes were not regulated by auranofin. Therefore, treatment solely with auranofin was insufficient for regulation of this module.

**BED inhibitors partially regulated the antioxidative stress response module.** Then, we asked whether any additional compounds or combinations thereof could collectively regulate the dysregulated “lightcyan” module as a whole, since it could not be regulated by auranofin. We sought to determine whether auranofin might be able to regulate the “lightcyan” module. Auranofin is used to treat rheumatoid arthritis. Auranofin is an inhibitor of thioredoxin reductase (encoded by TXNRD1/2)32. This drug is used as an approved drug to treat rheumatoid arthritis.

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We analyzed the transcriptome and epigenome perturbation datasets of the compounds with desirable profiles and found that the BET inhibitors (+)-JQ1 and GSK1210151A may be candidates that could downregulate the expression of group 1 genes (Fig. 5a). Group 1 included genes downregulated by BET inhibitors, such as GPX2 and ALDH3A1. On the other hand, genes upregulated by BET inhibitors were also detected in a different group, designated as group 2, which included IDH1 and GSR.

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downregulated by BET inhibitors, which decreased the chromatin accessibility of their promoters or enhancers in a concentration-dependent manner (Figs. 5b and S10). We further investigated whether sensitivity to auranofin could be changed through transcriptional perturbations induced by BET inhibitors. We treated the cell lines with BET inhibitors combined with auranofin. In H2228 cells, which showed one of the highest activity levels in terms of the “lightcyan” module, auranofin and (+)-JQ1 showed synergistic activity (CI50 = 0.56) (Fig. 5c).

Despite the successful combined use of auranofin and (+)-JQ1, we found that the combination did not exert significant synergistic effects on some cell lines, such as H2126, II-18 and other cell lines (Figs. 5c and S11). We found that the combination of these compounds did not regulate the expression of group 2 genes (Fig. 5a).

We additionally examined whether there were any other epigenetic drugs that could specifically control the expression of group 2 genes based on the observed omics information. As a candidate in place of auranofin, we used vorinostat with (+)-JQ1 treatment to more widely and moderately modulate the “lightcyan” module (Supplementary Fig. S12), and we observed a slight synergistic effect for (+)-JQ1 and vorinostat in some cells (CI40 = 0.61 in H2228 cells and CI40 = 0.80 in H2126 cells) with high activity in the “lightcyan” module (Fig. 5d).
To further elucidate the possible mode of action, we scrutinized the differences in the epigenome status between genes controlled and uncontrolled by BET inhibitors. For example, NFE2L2 binding sites reside in the enhancer region of ALDH3A1, which is one of the controlled genes. In this case, the chromatin structures represented in the ATAC-seq data were closed, and transcriptional activity was decreased by the BET inhibitors. In contrast, in GSR, the strong binding of NFE2L2 was localized to the promoter region, and the BET inhibitor did not close the associated chromatin structure, although a previously unknown enhancer located in the gene body (Fig. 5b) was inaccessible. These findings suggest that the regulatory mechanisms of NFE2L2 binding may be promoter- and enhancer-specific. In another example, OSGIN1 was significantly upregulated in response to BET inhibitors (Figs. 5a and S10). OSGIN1 is an oxidative stress response gene that is associated with the regulation of apoptosis. A regulatory region resides 25 kb upstream of this gene. We found that the chromatin structure was closed in response to treatment with BET inhibitors. When we analyzed the ENCODE ChIP-seq data, we found that this region overlapped with the binding site of the CTCF/cohesin complex. The BET inhibitors may have changed the accessibility of the insulator regions, which may have affected the chromatin structure and transcriptional activity. We suggest that improving the understanding of the transcriptional and epigenomic characteristics might help us to completely regulate the module.

Finally, we named the “lightcyan” module the “redox” module which was associated with KEAP1-NFE2L2 dysregulation, metabolic features of antioxidative pathways and auraofin sensitivity and could be partly regulated by epigenetic inhibitors.

### Possible cell type-specific vulnerable modules.

Overall, the above results indicate that the possible vulnerable modules, which should serve as the most effective targets for drug interventions, may differ depending on the cell type. Even when the putative redox module is not fully regulated in some cell lines, other modules may be targeted by other drug combinations. To further address this issue, we reorganized and characterized each of the lung cancer cell lines according to the patterns of the modules. We found that certain modules were associated with the lineages and differentiated phenotypes of the cells. We identified five modules that included representative markers of the cellular lineages and differentiated phenotypes. For example, the “brown” module included genes encoding epithelial markers such as E-cadherin (CDH1), EPCAM, GATA6 and CPA. The “violet” module included the alveolar type-I and -II cell markers HOPX and AQP5 and the surfactant protein A genes (SFTPA1, SFTPA2, and SFTA2). The “violet” module also contained genes associated with the inflammatory response.

In contrast, the “midnightblue” module, which showed a significantly negative correlation with the “brown” module (r = −0.89, p = 4.0e-10, Pearson correlation coefficient) included the epithelial-mesenchymal transition regulator ZEB1. The “purple” and “blue” modules included bronchial markers of basal and neuroendocrine cells. Based on the activity patterns of these characteristic modules, we clustered the cell lines depending on their omics signatures (Fig. 6a).

Unlike classifications based on representative surface markers and genomic mutations, this module-based classification should reflect system-level cancerous aberrations. Indeed, when we mapped the activities of the modules to the clinical samples, the clinical samples of lung cancers exhibited diverse module characteristics (Fig. 6b). For example, the patterns of module activity were occasionally associated with the degree of malignancy, as indicated by the presence of poor prognosis (Fig. 6c). Moreover, in vivo cancers frequently showed remarkably diverse variations, although their driver mutations were the same. Distinct therapeutic strategies may be required depending on the unique multi-omic and phenotypic characteristics of different cancer types.

To develop a new drug for each of these cancer subtypes, appropriate cell lines should be selected as models. It follows that the reclassification of cancers according to their gene expression modulations, in addition to the use of the current simple classification scheme based on their genomic mutational patterns, should provide clues regarding the mechanisms of oncogenesis and tumor progression that could be pharmacologically targeted.

### Methods

#### Cell lines.

Human lung cancer cell lines were cultured and harvested as previously described. They are described in our database DBKERO (https://kero.hgc.jp/) for high-throughput RNA-seq and ATAC-seq. Cells (1,500–12,000 cells per well) were seeded and treated with compounds in 96-well culture plates and then cultured for 24–72 h.

#### Chemical compounds.

Cells were treated with 320 compounds in the viability test. For this purpose, we selected 305 compounds from the Library of Pharmacologically Active Compounds (LOPAC, Sigma-Aldrich), 10 epigenetically targeted drugs and 4 tyrosine kinase inhibitors.
We designed high-throughput RNA-seq and ATAC-seq analyses that utilized drug treatment in a 96-well plate format. Two types of drug plates were prepared: dataset 1 included 24 compounds (23 compounds + a DMSO control) at four concentration points, and dataset 2 included 96 compounds (95 compounds + a DMSO control) at a single concentration and time point. For the RNA-seq and ATAC-seq analyses, 23 and 95 compounds, respectively, were selected from among the same compounds used for the viability test. According to their functional categories and growth inhibition rates, the compound showing some changes on cellular phenotypes were mainly selected. Some compounds were additionally selected from the FDA-approved Drug Library (Selleck Chemicals and Prestwick Chemical). The complete list is shown in Supplementary Table S1.

**High-throughput RNA-seq.** The culture medium was discarded from a 96-well culture plate kept on ice. The cells were washed using cold PBS. Total RNA was extracted using TRIZol (Invitrogen). Briefly, 100 μl TRIZol was added to each well, and the cells were lysed. We prepared a new 1.5 ml tube with 400 μl TRIZol and transferred the lysed cells into the tube. For RNA separation, 100 μl chloroform was added and mixed well, and the sample was incubated at room temperature for 10 min. After the incubation, the sample was centrifuged (12000 rpm, 15 min, 4°C), and the supernatant was transferred to a new 1.5 ml tube. Isopropanol precipitation was performed for RNA purification. The RNA sample was eluted in 10 μl nuclease-free water.

To perform the automatic reverse transcription and amplification reactions in microchambers, we used the C1 Single-Cell Auto Prep System (Fluidigm) with a modified script (provided at https://kero.hgc.jp/). The detailed procedure is shown in the Supplementary Methods. Briefly, reverse transcription and cDNA amplification were performed using the C1 Integrated Fluidic Circuit (C1 Single-Cell Auto Prep IFC for Open App, Fluidigm) with the C1 system. The purified cDNAs were used for library construction with the Nextera XT DNA Sample Preparation Kit (Illumina). The library was sequenced with a 35-base single-end run using the HiSeq. 2500 System (Illumina).

**High-throughput ATAC-seq.** We developed a 96-well format ATAC-seq procedure according to the original ATAC-seq protocol with some modifications. The detailed procedure is shown in the Supplementary Methods. Briefly, the cells were dissociated using 0.25% Trypsin-EDTA (25200–056, Gibco) or AccutaseTM (A11105-01, Gibco) and pipetted into 8-channel tubes. For cell lysis, 50 μl lysis buffer (10 mM Tris-HCl pH 7.4, 10 mM NaCl, 3 mM MgCl₂, and 0.1% NP-40) was used. For the ATAC reaction, 10 μl of 2 × TD buffer, 1 μl of Tn5 transposase and 4 μl of nuclease-free water were added, and the sample was incubated at 37 °C for 30 min. The transposed DNAs were purified using the ZR-96 DNA Clean & Concentrator-5 (Zymo Research) and eluted with 25 μl of nuclease-free water. PCR amplification was performed using 19.7 μl of transposed DNAs according to the original ATAC-seq protocol, and then 10 μl of the sample in each of the 96 wells was mixed and purified. The libraries were sequenced with a 35-base single-end run using the HiSeq. 2500 platform (Illumina).

**ATAC-seq under basal conditions.** To evaluate the open chromatin patterns of lung cancer cell lines under basal conditions, approximately 5 × 10⁴ cells were prepared and treated according to the ATAC-seq protocol previously described.

**Determination of the promoter and enhancer regions.** To determine the promoter regions, we selected the representative transcriptional start sites (TSSs) according to the start positions of the RefSeq transcripts of each gene using TSS-seq data (accession number DRA005903). We determined 13,431 promoter regions located within ±1.5 kb from representative TSSs.

To define the enhancer regions, H3K27ac and H3K4me1 ChIP-seq data from 27 cell lines were used (accession numbers DRA001860 and DRA002311). The MACS2 peaks from the 27 cell lines were merged and the promoter regions were removed using bedtools. We identified 312,226 and 535,538 regions according to the H3K27ac and H3K4me1 ChIP-seq peaks, respectively. All regions were merged using bedtools, and 515,599 regions were identified as enhancer regions. We used 56,961 enhancer regions that overlapped the ATAC-seq peaks under basal conditions. We assigned 32,111 enhancer regions to genes (±100 kb from TSSs).

**Analysis of the high-throughput RNA-seq and ATAC-seq data.** RNA-seq tags were mapped to the reference genome (UCSC hg19) using ELAND software (Illumina). The tags were mapped to the sequences obtained from the three RNA spike in controls. The rpkm values were calculated for 22,657 genes and the spike-in reference genome (UCSC hg19) using ELAND software (Illumina). The tags were mapped to the sequences obtained from the three RNA spike in controls. The rpkm values were calculated for 22,657 genes and the spike-in reference genome (UCSC hg19) using ELAND software (Illumina). The peaks were detected using MACS2 with the default parameters. ATAC-seq data with >0.5 million mapped reads, 2) <15% intron mapped reads, and 3) rpkm values for the spike-in control that were within ±2 sd of the average for each plate. For the analysis of reproducibility, we used protein-coding genes with rpkm values ≥2 in both datasets. The ATAC-seq tags were mapped to the reference genome (UCSC hg19) using ELAND software (Illumina). The peaks were detected using MACS2 with the default parameters. ATAC-seq data with >10,000 MACS peaks were used for further analysis. The ppb values were calculated for the promoter and enhancer regions. As ELAND software is a somewhat older software, we compared the mapping results from ELAND with those from other tools and confirmed that there were no significant differences among the results of the mapping tools for the analysis of the high-throughput RNA-seq and ATAC-seq data (Supplementary Fig. S3d,g).

**CCLE RNA-seq data.** For the lung adenocarcinoma cell lines A549 and H1299, RNA-seq data (CCLE_ RNAseq_081117.rpkm.gct) were downloaded from the CCLE download page (https://portals.broadinstitute.org/ccle/data). The rpkm values were log2-tranformed after +1. For the comparison of the datasets obtained in this study, we used genes with rpkm values ≥2 in both datasets.
qPCR validation of RNA-seq and ATAC-seq data. For the cDNAs that were used in the high-throughput RNA-seq analysis, qPCR validation was performed using THUNDERBIRD SYBR qPCR Mix (QPS-201, TOYOBO). We performed one assay for each measurement. The detailed procedure is shown in the Supplementary Methods. The primer sequences are listed in Supplementary Table S6.

We also performed qPCR analyses of the five promoters and eight enhancers for the validation of the high-throughput ATAC-seq data. We performed one qPCR assay for each measurement. The ATAC DNAs, which were subjected to sequence analysis, were used. The detailed procedure is shown in the Supplementary Methods. The primer sequences are listed in Supplementary Table S7.

Cell viability test. Viability tests of cells treated with 320 drugs were performed using a 384-well plate high-throughput system. Each well of a 384-well plate contained 7.5 μl of culture medium. DMSO control wells were prepared with or without cells to serve as positive and negative controls, respectively. For high-throughput stimulation, 2.5 μl of a chemical compound was added to each well of the plates. For each well, 450–3,000 cells (15 μl of a cell suspension) were seeded and incubated for 72 h at 37 °C in an atmosphere containing 5% CO₂. After incubation, 25 μl of the reagent provided with the CellTiter-Glo Luminescent Cell Viability Assay Kit (Promega) was added. A Multidrop Combi (Thermo Fisher Scientific) and a HORNET-NX (Wako) automatically dispensed the assay components. The cell viability rates were calculated according to the luminescence, which was measured using an EnVision Plate Reader (PerkinElmer). The assays were repeated twice (N = 2). The data were fitted to the logistic equation using XLfit (IDBS) as follows:

\[
y = A + \frac{B - A}{1 + \left(\frac{x}{C}\right)^D}
\]

The values represented the variables as follows: y: % growth inhibition; x: concentration of the compound; A: minimum value of y; B: maximum value of y; C: EC50 value; D: Hill slope.42

Analysis of the association between the basal genomic status and drug sensitivity. The 26 cell lines were divided into two groups according to the IC₅₀ values of each compound to produce the maximum degree of separation in terms of the IC₅₀ values between the groups (Fig. 2c). The IC₅₀ values were adjusted to 10 μM when the growth inhibition rate did not reach 50% at a 10 μM concentration. We focused on the mutational status of 26 cancer-related genes and used Fisher’s exact test to analyze the association between the mutational status and the cell groups defined by the IC₅₀ value. An aberrant status was defined according to the following: for oncogenes, mutants were defined if the cell lines harbored genomic mutations and the expression was >5 rpm; for tumor-suppressor genes, mutants were defined if they harbored mutations or showed low expression levels that were <5 rpm. For other cancer-related genes, such as those encoding chromatin factors and fusion genes, mutants were defined if they showed genomic mutations. For CDKN2A, the DNA methylation status of the p16INK4A alternative promoter was considered. Finally, we extracted the gene–compound pairs with a >5-fold IC₅₀ differences between the two cell groups and a significant enrichment of the mutational status of the gene in either group (p < 0.05, Fisher's exact test).

CMAP analysis. Using the high-throughput RNA-seq data, genes with >4 or <0.25-fold changes in terms of their expression levels were extracted as parts of the signatures of the transcriptome mode of action. We used an RNA-seq dataset from PC-9 cells treated with vorinostat (0.95 μM for 24 h). Of the 4,798 genes expressed in either stimulated or DMSO control cells (rpm value > 10 for RNA-seq; ppm > 5 ppm from the promoter region for ATAC-seq), 163 upregulated and 146 downregulated genes were extracted and designated a part of the signature of our dataset. To verify the signature, we accessed CMAP datasets using CLUE. We used the CLUE L1000 Query to evaluate the similarities between the signature of our dataset to that of the CMAP datasets.

Weighted gene co-expression network. For the extraction of the gene co-expression modules, we used RNA-seq data (accession numbers DRA001846 and DRA002311) from 27 cultured cell lines (26 lung cancer cell lines and small airway epithelial cells) incubated under basal conditions. We extracted 73 modules using the WGCNA method.45 The detailed procedure is shown in the Supplementary Methods. The complete list of the 73 module members is included in Supplementary Table S3. GO enrichment analysis of each module was performed using the WGCNA GOenrichmentAnalysis function. The top 10 GO terms ordered according to the Bonferroni corrected p-values (Bonferroni p < 0.1) are listed in Supplementary Table S4.

Analysis of the NFE2L2 binding patterns using ENCODE data. To evaluate the NFE2L2 binding patterns, NFE2L2 ChIP-seq data (ENCs584GHV and NCSR949BZP) were obtained from the ENCODE project.41 The fastq files (ENCFF002EAK and ENCF002ECM) were mapped to UCSC hg19 using Bowtie2 (v2.0.6).46 The sorted bam files were created using SAMTools.47 The peaks were detected using MACS2 (v2.1.0.20150420).

Metabolome analysis. For the metabolome analyses, capillary electrophoresis time-of-flight mass spectrometry (CE-TOFMS) was performed using an Agilent CE-TOFMS system. The raw data were analyzed using MasterHands (ver. 2.17.0.18)48. For further analysis, the average concentrations (fmol/cell, n = 2) were used for each cell line. A not detected (N.D.) value was omitted if the compound was detected using one measurement. Concentrations were considered as zero if the metabolites were not detected by either measurement.

To evaluate the association with the transcriptome modules, Spearman correlation coefficients for the ME values of the modules and the concentrations of the metabolites were calculated. We detected 113 metabolites in >10 cell lines.
siRNA knockdown experiments. A549 cells were seeded (1500 cells/well in 96-well plates) and incubated for 24h at 37°C. NFE2L2 siRNAs (Silencer Select NFE2L2 s9491 and s9493, Applied Biosystems) and negative control siRNAs (Silencer Select Negative Control #1 siRNA, Applied Biosystems) were used for the transfection of the A549 cells. Lipofectamine RNAiMAX Transfection Reagent (Thermo Fisher Scientific) and the siRNAs were diluted in serum-free MEM (M7278, Sigma-Aldrich). For the preparation of the RNA lipid complexes, diluted Lipofectamine and siRNAs were mixed and incubated for 5 min, and 10μl of siRNA-lipid reagents was added to the cells.

To confirm the knockdown efficiency and determine the changes in the expression levels of the downstream genes, qRT-PCR was conducted (n = 3). For qRT-PCR, cDNAs were synthesized using the SuperPrep Cell Lysis & RT Kit for qPCR (SQ-101, TOYOBO), and qPCR was performed using the THUNDERBIRD SYBR qPCR Mix (QPS-201, TOYOBO). The PCR primers are listed in Supplementary Table S8.

Analysis of TCGA data. For the data from TCGA for lung adenocarcinoma (LUAD), lung squamous cell carcinoma (LUSC) and esophageal carcinoma (ESCA), RNA-seq v2 data was acquired using TCGA-Assembler v2.0.6 (the data checked on 2019/09/10). The clinical information was acquired from cBioPortal (downloaded on 2019/09/13). The somatic mutation data were downloaded from the NCI Genomic Data Commons (https://gdc.cancer.gov/) (downloaded on 2019/09/13–14). To generate the heat maps of the RNA-seq v2 data, the normalized counts were adjusted to 1 if the count was <1. The adjusted expression levels were normalized to the median expression levels in all patients and log2-transformed. Hierarchical clustering with Ward’s method was performed according to the Euclidean distance using the expression levels of the genes in the given modules.

To extract the core putative redox module member genes (Fig 5a) that exhibited significant co-expression patterns in the clinical samples, we performed the Wilcoxon rank-sum test to compare the expression levels of TCGA-LUAD cases with high module activity (Fig. 4d) with those of other cases. We extracted 63 differentially co-expressed genes among the high module activity cases (Bonferroni p < 0.05).

To associate the TCGA transcriptome data with the modules, the “module score” for each TCGA-LUAD case was calculated for each module as follows: (1) the expression values (normalized counts) were log2-transformed after adding 1; (2) the log2-transformed values of 10 representative genes with the highest correlation with the ME were averaged for each case; (3) z-score normalization was applied. Kaplan-Meier analysis and the log-rank test were performed using the survival package in R.

Treatment with a combination of auranofin and (+)-JQ1. For the combination treatment using auranofin and (+)-JQ1, the assay plate was designed as shown in Supplementary Fig. S11. The cell viability rates were calculated as described above. The combination index (CI) was calculated according to the Chou-Talalay method48.

Treatment with a combination of vorinostat and (+)-JQ1. For the combination treatment using vorinostat and (+)-JQ1, the CI was calculated in a manner similar to that shown in Supplementary Fig. S11. By using three cell lines with high activity in the putative redox module, a cell viability test was performed twice, and the CI was determined according to the average cell viability rates.

Data availability

All the raw sequencing data used for the high-throughput RNA-seq and ATAC-seq analyses have been registered in DDBJ under the accession numbers DRA006875 - DRA006902 (RNA-seq) and DRA006903 - DRA006930 (ATAC-seq). The omics sequencing data for the cell lines were previously published48,49. The datasets in this paper are provided in our database, DBKERO (https://kero.hgc.jp/)22.

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Additional information

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