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# Comprehensive genomic characterization of squamous cell lung cancers

The Cancer Genome Atlas Research Network

## **Summary**

Lung squamous cell carcinoma (lung SqCC) is a common type of lung cancer, causing approximately 400,000 deaths per year worldwide. Genomic alterations in lung SqCC have not been comprehensively characterized and no molecularly targeted agents have been developed specifically for its treatment. As part of The Cancer Genome Atlas (TCGA), we profiled 178 lung SqCCs to provide a comprehensive landscape of genomic and epigenomic alterations. Lung SqCC is characterized by complex genomic alterations, with a mean of 360 exonic mutations, 165 genomic rearrangements, and 323 segments of copy number alteration per tumor. We found statistically recurrent mutations in 18 genes in including mutation of *TP53* in nearly all specimens. Previously unreported loss-of-function mutations were seen in the *HLA-A* class I major histocompatibility gene. Significantly altered pathways included *NFE2L2/KEAP1* in 34%, squamous differentiation genes in 44%, *PI3K/AKT* in 47%, and *CDKN2A/RB1* in 72% of tumors. We identified a potential therapeutic target in the majority of tumors, offering new avenues of investigation for lung SqCC treatment.

#### Introduction

Lung cancer is the leading cause of cancer-related mortality worldwide, leading to an estimated 1.4 million deaths in  $2010^1$ . The discovery of recurrent mutations in the Epidermal Growth Factor Receptor (*EGFR*) kinase as well as fusions involving the Anaplastic Lymphoma Kinase (*ALK*), has led to a dramatic change in the treatment of

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We specifically recognize the following investigators who made substantial contributions to the project.

Manuscript co-ordinators: P.S.H. and D.N.H. Data co-ordinator: M.D.W. Analysis co-ordinators: P.S.H. and N.S. DNA sequence analysis: P.S.H., M.S.L., A.S., B.H. and G.G. mRNA sequence analysis: M.D.W., J.L. and D.N.H. DNA methylation analysis: L.C., J.G.H. and L.D. Copy number analysis: A.C., G.S., N.H.P., R.K. and M.L. Pathway analysis: N.S., R.B., C.J.C., R.S., C.M., S.N., E.A.C., R.S., J.N.W. and C.S. miRNA sequence analysis: A.C. and G.R. Pathology and clinical expertise: W.D.T., B.E.J., D.A.W. and M.S.T. Project chairs: S.B.B., R.G. and M.M.

**Author Information:** The primary and processed data used to generate the analyses presented here can be downloaded by registered users at https://tcga-data.nci.nih.gov/tcga/tcgaDownload.jsp, https://cghub.ucsc.edu/ and https://tcga-data.nci.nih.gov/docs/publications/lusc\_2012/

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patients with lung adenocarcinoma, the most common type of lung cancer<sup>2–5</sup>. More recent data have suggested that targeting mutations in *BRAF*, *AKT1*, *ERBB2* and *PIK3CA* and fusions that involve *ROS1* and *RET* may also be successful<sup>6,7</sup>. Unfortunately, activating mutations in *EGFR* and *ALK* fusions are typically not present in the second most common type of lung cancer, lung squamous cell carcinoma (lung SqCC),<sup>8</sup> and targeted agents developed for lung adenocarcinoma are largely ineffective against lung SqCC.

Although no comprehensive genomic analysis of lung SqCCs has been reported, single-platform studies have identified regions of somatic copy number alteration in lung SqCCs, including amplification of *SOX2*, *PDGFRA* and *FGFR1/WHSC1L1* and deletion of *CDKN2A*<sup>9,10</sup>. DNA sequencing studies of lung SqCCs have reported recurrent mutations in a number of genes, including *TP53*, *NFE2L2*, *KEAP1*, *BAI3*, *FBXW7*, *GRM8*, *MUC16*, *RUNX1T1*, *STK11* and *ERBB4*<sup>11,12</sup>. *DDR2* mutations and *FGFR1* amplification have been nominated as therapeutic targets <sup>13–15</sup>.

We have conducted a comprehensive study of lung SqCCs from a large cohort of patients as part of The Cancer Genome Atlas (TCGA) project. The twin aims are to characterize the genomic/epigenomic landscape of lung SqCC and to identify potential opportunities for therapy. We report an integrated analysis based on DNA copy number, somatic exonic mutations, mRNA sequencing (RNA-seq), mRNA expression, and promoter methylation for 178 histopathologically reviewed lung SqCCs in addition to whole-genome sequencing (WGS) of 19 samples and miRNA sequencing of 159 samples (Supplementary Table S1.1). Demographic and clinical data and results of the genomic analyses can be downloaded from the TCGA data portal (https://tcga-data.nci.nih.gov/tcga/tcgaDownload.jsp).

## Samples and clinical data

Tumor samples were obtained from 178 patients with previously untreated stage I–IV lung SqCC. Germline DNA was obtained from adjacent, histologically normal tissues resected at the time of surgery (n=137) or peripheral blood (n=41). All patients provided written informed consent to conduct genomic studies in accordance with local Institutional Review Boards. The demographic characteristics are described in Supplementary Table S1.2. The median follow-up for the cohort was 15.8 months, and 60% of patients were alive at the time of last follow-up (data updated in November 2011). 96% of the patients had a history of tobacco use, similar to previous reports for North American lung SqCC patients <sup>16</sup>. DNA and RNA were extracted from patient specimens and were measured by several genomic assays, which included standard quality control assessments (Supplementary Methods, sections 2–8). A committee of experts in lung cancer pathology performed an additional review of all samples to confirm the histological subtype (Supplementary Figure 1.1 and Supplementary Methods, section 1).

#### **Somatic DNA Alterations**

The lung SqCCs analyzed in this study display a large number and variety of DNA alterations, with a mean 360 exonic mutations, 323 altered copy number segments and 165 genomic rearrangements per tumor.

Copy number alterations were analyzed using multiple platforms. Analysis of SNP 6.0 array data across the set of 178 lung SqCCs identified a high rate of copy number alteration (mean of 323 segments) when compared to other TCGA projects (as of February 1, 2012) including ovarian cancer (477 segments), <sup>17</sup> glioblastoma multiforme (282 segments), <sup>18</sup> colorectal carcinoma (213 segments), breast carcinoma (282 segments) and renal cell carcinoma (156 segments). These segments gave rise to regions of both focal and broad somatic copy number alterations (SCNAs) with a mean of 47 focal and 23 broad events per tumor (broad events defined as 50% of the length of the chromosome arm). There was strong concordance between the three independent copy number assays for all regions of SCNA (Supplementary Figures S2.1–2.4).

At the level of whole chromosome arm SCNAs, lung SqCCs exhibit many similarities to 205 cases of lung adenocarcinoma analyzed by TCGA (Supplementary Figure S2.1a). The most striking difference between these cancers is selective amplification of chromosome 3q in lung SqCC, as has been reported<sup>9,19</sup> (p<1×10<sup>-15</sup> by Fisher's exact test). Using the SNP 6.0 array platform and GISTIC 2.0<sup>20,21</sup>, we identified regions of significant copy number alteration (Supplementary Methods, section 2). There were 50 peaks of significant amplification or deletion (q<0.05), several of which included SCNAs previously seen in lung SqCCs including *SOX2*, *PDGFRA/KIT*, *EGFR*, *FGFR1/WHSC1L1*, *CCND1* and *CDKN2A*<sup>9,10,19</sup> (Supplementary Figure S2.1b and Supplementary Data S2.1 and S2.2). Other peaks defined regions of SCNA reported for the first time including amplifications of chromosomal segments containing *NFE2L2*, *MYC*, *CDK6*, *MDM2*, *BCL2L1* and *EYS* and deletions of *FOXP1*, *PTEN* and *NF1* (Supplementary Figure S2.1b).

Whole exome sequencing of 178 lung SqCCs and matched germline DNA, targeted 193,094 exons from 18,863 genes. Mean sequencing coverage across targeted bases in was 121X with 83% of target bases above 30X coverage. We identified a total of 48,690 non-silent mutations with a mean of 228 non-silent and 360 total exonic mutations per tumor, corresponding to a mean somatic mutation rate of 8.1 mutations per megabase (Mb) and median of 8.4/Mb. That rate is higher than rates observed in other TCGA projects including acute myelogenous leukemia (0.56/Mb), breast carcinoma (1.0/Mb), ovarian cancer<sup>17</sup> (2.1/Mb), glioblastoma multiforme<sup>18</sup> (2.3/Mb) and colorectal carcinoma (3.2/Mb) (data as of February 1, 2012, p<2.2 x10<sup>-16</sup> by t-test or Wilcoxon rank sum test for lung SqCC versus all others). In lung SqCC, CpG transitions and transversions were the most commonly observed mutation types, with mean rates of 9.9 and 10.7 per sequenced megabase of CpG context, respectively, for a total mutation rate of 20.6/Mb. At non-CpG sites, transversions at C:G sites were more common than transitions (7.3 vs. 2.9 per Mb; total = 10.2/Mb) and more common than transversions or transitions at A:T sites (1.5 vs. 1.3 per Mb; total = 2.8/Mb).

Significantly mutated genes were identified using a modified version of the MutSig algorithm (Supplementary Methods, section 3)<sup>22,23</sup>. We identified 10 genes with a false discovery rate (FDR) q value <0.1 (Supplementary Table S3.1): *TP53, CDKN2A, PTEN, PIK3CA, KEAP1, MLL2, HLA-A, NFE2L2, NOTCH1* and *RB1*, all of which demonstrated robust evidence of gene expression as defined by Reads Per Kilobase of exon model per Million mapped reads (RPKM) >1 (Figure 1). *TP53* mutation was observed in 81% of samples by automated analysis; visual review of sequencing reads identified an additional

9% of samples with potential mutations in regions of sub-optimal coverage or in samples with low purity. The majority of observed mutations in *NOTCH1* (8 of 17) were truncating alterations, suggesting loss-of-function, as has recently been reported for head and neck SCCs<sup>22,24</sup>. Mutations in *HLA-A* were also almost exclusively nonsense or splice site events (7 of 8).

To increase our statistical power to detect mutated genes in the setting of the observed high background mutation rate, we performed a secondary MutSig analysis only considering genes previously observed to be mutated in cancer according to the COSMIC database. This yielded 12 additional genes with FDR <0.1: FAM123B (WTX), HRAS, FBXW7, SMARCA4, NF1, SMAD4, EGFR, APC, TSC1, BRAF, TNFAIP3 and CREBBP (Supplementary Table S3.1). Both the spectrum and the frequency of EGFR mutations differed from those seen in lung adenocarcinomas. The two most common alterations in lung adenocarcinoma, L858R and in-frame deletions in exon 19, were absent, whereas two L861Q mutations were detected in EGFR.

As described in Supplemental Figure S3.1 we verified somatic mutations by performing an independent hybrid-recapture of 76 genes in all samples. 1289 mutations were assayed and we achieved satisfactory coverage to have power to verify at 1283 positions. 1235 mutations were validated (96.2%) (Supplementary Figure S3.1 and Supplementary Methods, section 3). We also verified mutation calls using WGS and RNA-sequencing data with similar results (Supplementary Figures S3.1 and S4.3 and Supplementary Methods, sections 3 and 4).

Whole genome sequencing was performed for 19 tumor/normal pairs with a mean computed coverage of 54X. A mean of 165 somatic rearrangements was found per lung SqCC tumor pair (Supplementary Figure S3.2), a value in excess of that reported for WGS studies of other tumor types including colorectal carcinoma (75)<sup>25</sup>, prostate carcinoma (108)<sup>26</sup>, multiple myeloma (21)<sup>23</sup>, and breast cancer (90)<sup>27</sup>. Although the majority of in-frame coding fusions detected in WGS were validated by RNA-seq, no recurrent rearrangements predicted to generate fusion proteins were identified (Supplementary Data S3.1 and S4.1).

## Somatically altered pathways

Many of the somatic alterations we have identified in lung SqCCs appear to be drivers of pathways important to the initiation or progression of the cancer. Specifically, genes involved in oxidative stress response and squamous differentiation were frequently altered by mutation or SCNA. We observed mutations and copy number alterations of *NFE2L2* and *KEAP1* and/or deletion or mutation of *CUL3* in 34% of cases (Figure 2). *NFE2L2* and *KEAP1* code for proteins that bind each other, have been shown to regulate the cell's response to oxidative damage, chemo- and radiotherapy and are somatically altered in a variety of cancer types<sup>28,29</sup>. We found mutations in *NFE2L2* almost exclusively in one of two KEAP1 interaction motifs, DLG or ETGE. Mutations in *KEAP1* and *CUL3* showed a pattern consistent with loss-of-function and were mutually exclusive with mutations in *NFE2L2* (Figures 1c and 2). PARADIGM<sup>30</sup> analysis predicts that mutations in *NFE2L2* and

*KEAP1* exert a significant functional impact (Supplementary Figure S7.C.1 and S.7.C.2 and Supplementary Methods, section 7).

We also found alterations in genes with known roles in squamous cell differentiation in 44% of samples, including overexpression and amplification of *SOX2* and *TP63*, loss-of-function mutations in *NOTCH1*, *NOTCH2*, and *ASCL4* and focal deletions in *FOXP1* (Figure 2). Although *NOTCH1* has been well characterized as an oncogene in hematologic cancers<sup>31</sup>, *NOTCH1* and *NOTCH2* truncating mutations have been reported in cutaneous SCCs and lung SqCCs<sup>32</sup>. Truncating mutations in *ASCL4* are the first to be reported in human cancer and may have a lineage role given the requirement for *ASCL1* for survival of small cell lung cancer cells<sup>33</sup>. Alterations in *NOTCH1*, *NOTCH2* and *ASCL4* were mutually exclusive and exhibited minimal overlap with amplification of *TP63* and/or *SOX2* (Figure 2), suggesting that aberrations in those modulators of squamous cell differentiation have overlapping functional consequences.

## mRNA expression profiling and subtype classification

Whole-transcriptome expression profiles were generated by RNA-sequencing for the entire cohort and by microarrays for a 121-sample subset. Of 20,502 genes analyzed, the mean RNA coverage indices were 19X and 6420 RPKM (Supplementary Figure S4.1 and Supplementary Methods, section 4). Previously reported lung SqCC gene expressionsubtype signatures<sup>34</sup> were applied to both of the expression platforms, yielding four subtypes designated as classical (36%), basal (25%), secretory (24%) and primitive (15%). The concordance of subtypes between the two platforms was high (94% agreement) (Supplementary Figure S4.2). Significant correlations were found between the expression subtypes and genomic alterations in copy number, mutation and methylation (Figure 3). The classical subtype was characterized by alterations in KEAP1, NFE2L2, and PTEN, as well as pronounced hypermethylation and chromosomal instability. The 3q26 amplicon was present in all of the subtypes, but it was most characteristic of the classical subtype, which also showed the greatest overexpression of three known oncogenes on 3q: SOX2, TP63, and PIK3CA. RNA sequencing data suggested that high expression levels of TP63, in samples with and without amplification of TP63, was associated with dominant expression of the deltaN isoform (also called p40), which lacks the N-terminal transactivation domain, compared to the longer isoform, called tap63 (89% of tumors overexpressed deltaN compared to tap63; p<2.2e-16). That short deltaN isoform is thought to function as an oncogene<sup>35,36</sup>, and its expression was most enriched in the classical subtype. In contrast, the primitive expression-subtype more commonly exhibited RB1 and PTEN alterations, and the basal expression-subtype showed NF1 alterations (Figure 3). Amplification of FGFR1 and WHSC1L1 was anti-correlated with the classical subtype and specifically with NFE2L2 or KEAP1 mutated samples. While CDKN2A alterations are common in lung SqCCs, they are not associated with any particular expression subtype (Figure 3).

Independent clustering of miRNA and methylation data indicated association with expression subtypes. The highest overall methylation was seen in the classical subtype (Figure 3, Supplementary Figures S5.1 and S6.1, Supplementary Methods, sections 5 and 6, Supplementary Data S6.1 and S6.2 and Supplementary Table 5.1). Integrative clustering

(iCluster)<sup>37</sup> of mRNA, miRNA, methylation, SCNA, and mutation data demonstrated concordance with the mRNA expression-subtypes and associated alterations. (Figure 3, Supplementary Figure S7. A.1 and Supplementary Methods, section 7). Independent correlation of somatic mutations, copy number alterations and gene expression signatures revealed significant subtype associations with alterations in the *TP53*, *PI3K*, *RB1* and *NFE2L2/KEAP1* pathways (Supplementary Figure S7. B.1 and Supplementary Methods, section 7).

## Integrated analysis of the tumor suppressor locus for CDKN2A

Integrated multi-platform analyses showed that *CDKN2A*, a known tumor suppressor gene in lung SqCC<sup>38</sup> that encodes the INK4A/p16 and ARF/p14 proteins, is inactivated in 72% of cases of lung SqCC (Figure 4a and Supplementary Data S7.1)—by epigenetic silencing by methylation (21%), inactivating mutation (18%), exon 1 $\beta$  skipping (4%), and homozygous deletion (29%).

Analysis of mRNA expression across the *CDKN2A* locus revealed four distinct patterns of *p16INK4* and *ARF* expression: complete absence of both *p16INK4* and *ARF* (33%); expression of high levels of both *p16INK4* and *ARF* (31%); high expression of *ARF* and absent *p16INK4* (31%); or expression of a transcript that represents a splicing of exon E1b from *ARF* with the shared E3 of *ARF* and *p16INK4*, generating a premature stop codon (4%) (Supplementary Figure S4.4). Almost all of the cases completely lacking *p16INK4* and *ARF* expression showed homozygous deletion (Figure 4b and Supplemental Data S7.1). In one case, *p16INK4* expression was detected but analysis of WGS data demonstrated an intergenic fusion event that resulted in detectable transcription between exon 1a of *p16INK4* and exon 18 of *KIAA1797* (Figures 4b, 4c). Interestingly, combined analysis of WGS and RNA-sequencing data identified tumor suppressor gene inactivation by intra- or interchromosomal rearrangement in *PTEN*, *NOTCH1*, *ARID1A*, *CTNNA2*, *VHL* and *NF1*, in 8 additional cases (Supplementary Data S3.1 and S4.1).

In addition to homozygous deletion, there are frequent mutational events in *CDKN2A* (Figure 4b and Supplementary Data 7.1). These account for 45% of the 56 cases with high *p16INK4* and *ARF* expression. Furthermore, methylation of the exon 1a promoter accounts for many other cases of *CDKN2A* inactivation (70% of lung SqCCs with *ARF* expression in the absence of detectable *p16INK4*). Seven additional tumors in the high-*ARF*/low-*INK4A* group had documented mutations of *INK4A*, primarily nonsense, suggesting nonsensemediated decay as a mechanism. Of the 28% of tumors without *CDKN2A* alterations, *RB1* mutations were identified in eight cases and *CDK6* amplification in one case (Figure 4d).

## Therapeutic targets

Molecularly targeted agents are now commonly used in patients with adenocarcinoma of the lung while no effective targeted agents have been developed specifically for lung SqCCs<sup>13</sup>. We analyzed our genomic data for evidence of the two common genomic alterations in adenocarcinomas of the lung: *EGFR* and *KRAS* mutation. Only one sample had a *KRAS* codon 61 mutation, and there were no exon 19 deletions or L858R mutations in *EGFR*.

However, amplifications of *EGFR* were found in 7% of cases as were two instances of the L861Q *EGFR* mutation, which confers sensitivity to erlotinib and gefitinib<sup>39</sup>.

The presence of new potential therapeutic targets in lung SqCC was suggested by the observation that 96% of tumors (171 of 178) contain one or more mutations in tyrosine kinases, serine/threonine kinases, PI3K catalytic and regulatory subunits, nuclear hormone receptors, G protein-coupled receptors, proteases and tyrosine phosphatases (Supplementary Figure S7.D.1a and Supplementary Data S7.2 and S7.3). 50–77% of the mutations were predicted to have a medium or high functional impact as determined by Mutation Assessor score<sup>40</sup> (Supplementary Figure S7.D.1a) and 39% of tyrosine and 42% of serine/threonine kinase mutations were located in the kinase domain. Many of the alterations were in known oncogenes and tumor suppressors, as defined in the COSMIC database (Supplementary Data S7.3).

We selected potential therapeutic targets based on several features, including (i) availability of an FDA-approved targeted therapeutic agent or one under study in current clinical trials (Supplementary Data S7.2) (ii) confirmation of the altered allele in RNA-sequencing; and (iii) Mutation Assessor score<sup>40</sup>. Using those criteria, we identified 114 cases with somatic alteration of a potentially targetable gene (64%) (Supplementary Figure S7.D.1b and Supplementary Data S7.4). Among these we identified three families of tyrosine kinases, the ERBBs, FGFRs and JAKs, all of which were found to be mutated and/or amplified<sup>41</sup>. As discussed for *EGFR*, the mutational spectra in these potential therapeutic targets differed from those in lung adenocarcinoma (Supplementary Figure S7.D.2).<sup>42</sup>

To complement a gene-centered search for potential therapeutic targets, we analyzed core cellular pathways known to represent potential therapeutic vulnerabilities: PI3K/AKT, RTK and RAS. Analysis of the 178 lung SqCCs revealed alteration in at least one of those pathways in 69% of samples after restriction of the analysis to mutations confirmed by RNA-sequencing and to amplifications associated with overexpression of the target gene (Figure 5). Mutational events that have been curated in COSMIC are additionally shown in Supplementary Figure S7D.2 as is the distribution of mutations, amplifications and overexpression of the genes depicted in Figure 5C. Specifically, one of three components of the PI3K/Akt pathway (PIK3CA, PTEN or AKT3) was altered in 47% of tumors and receptor-tyrosine kinase (RTK) signaling likely altered by EGFR amplification, BRAF mutation or FGFR amplification or mutation in 26% of tumors. (Figures 5 and Supplementary Figure S7.D.3). Alterations in the PI3K/Akt pathway genes were mutually exclusive with EGFR alterations as identified by MEMo<sup>43</sup> (Supplementary Figure S7.D.4.). While the dependence of lung SqCC on many of these individual alterations remains to be defined functionally, this analysis suggests new areas for potential therapeutic development in this cancer.

#### **Discussion**

Lung SqCCs are characterized by a high overall mutation rate of 8.1 mutations/Mb and marked genomic complexity. Similar to high-grade serous ovarian carcinoma<sup>17</sup>, almost all lung SqCCs display somatic mutation of *TP53*. There were also frequent alterations in the

CDKN2A/RB1, NFE2L2/KEAP1/CUL3, PI3K/AKT and SOX2/TP63/NOTCH1 pathways, providing evidence of common dysfunction in cell cycle control, response to oxidative stress, apoptotic signaling and/or squamous cell differentiation. Pathway alterations clustered according to expression-subtype in many cases, suggesting that those subtypes have a biological basis.

A role for somatic mutation in the cancer hallmark of "avoiding immune destruction<sup>44</sup> is suggested by the presence of inactivating mutations in the *HLA-A* gene. Somatic loss-of-function alterations of *HLA-A* have not been reported previously in genomic studies of lung cancer. Given the recently reported efficacy of anti-Programmed Death 1 (PD1)<sup>45</sup> and anti-Cytotoxic T-Lymphocyte Antigen 4 (CTLA4) antibodies in NSCLC<sup>46</sup>, these *HLA-A* mutations suggest a possible role for genotypic selection of patients for immunotherapies.

Targeted kinase inhibitors have been successfully used for treatment of lung adenocarcinoma but minimally so in lung SqCC. The observations reported here suggest that a detailed understanding of the possible targets in lung SqCCs may identify targeted therapeutic approaches. While *EGFR* and *KRAS* mutations, the two most common oncogenic aberrations in lung adenocarcinoma, are extremely rare in lung SqCC, alterations in the FGFR kinase family are common in lung SqCC. Lung SqCCs also share many alterations in common with head and neck squamous cell carcinomas without evidence of human papilloma virus (HPV) infection, including mutation in *PIK3CA*, *PTEN*, *TP53*, *CDKN2A*, *NOTCH1* and *HRAS*<sup>22,24</sup>, suggesting that the biology of these two diseases may be similar.

The current study has identified a potentially targetable gene or pathway alteration in the majority of the lung SqCC samples studied. The data presented here can help organize efforts to analyze lung SqCC clinical tumor specimens for a panel of specific, actionable mutations to select patients for appropriately targeted clinical trials. These data could thereby help to facilitate effective personalized therapy for this deadly disease.

## **Methods Summary**

All specimens were obtained from patients with appropriate consent from the relevant institutional review board. DNA and RNA were collected from samples using the Allprep kit (Qiagen). We used commercial technology for capture and sequencing of exomes from tumor DNA and normal DNA and whole-genome shotgun sequencing. Significantly mutated genes were identified by comparing them with expectation models based on the exact measured rates of specific sequence lesions. GISTIC<sup>23,24</sup> analysis of the circular-binary-segmented Affymetrix SNP 6.0 copy number data was used to identify recurrent amplification and deletion peaks. Consensus clustering approaches were used to analyze mRNA, miRNA and methylation subtypes using previous approaches<sup>20,21,34,38,41,44</sup>.

## **Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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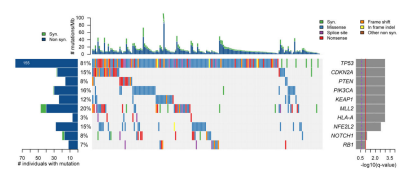


Figure 1. Significantly mutated genes in lung SqCC

Significantly mutated genes (q-value <0.1) identified by exome sequencing are listed vertically by q-value. The percentage of lung SqCC samples with a mutation detected by automated calling is noted at the left. Samples displayed as columns, with the overall number of mutations plotted at the top and samples arranged to emphasize mutual exclusivity among mutations.

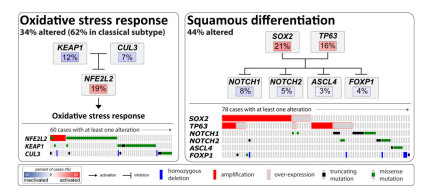


Figure 2. Somatically altered pathways in squamous cell lung cancer

**Left,** Alterations in oxidative response pathway genes by somatic mutation as defined by somatic mutation, copy number alteration or up- or down-regulation. Frequencies of alteration are expressed as a percentage of all cases, with background in red for activated genes and blue for inactivated genes. **Right,** Alterations in genes that regulate squamous differentiation, as defined in the left panel.

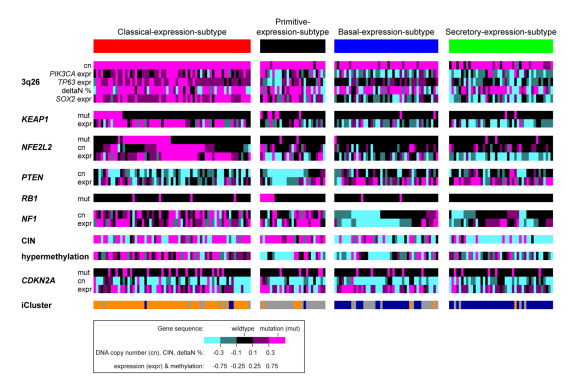
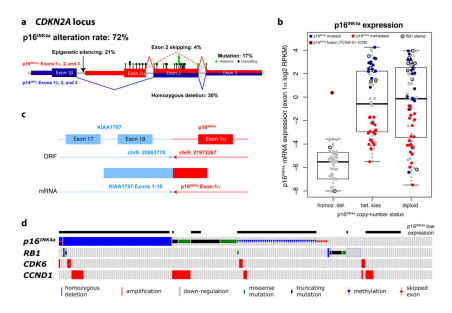


Figure 3. Gene expression subtypes integrated with genomic alterations

Tumors are displayed as columns, grouped by gene expression subtype. Subtypes were compared by Kruskal-Wallis tests for continuous features and by Fisher's exact tests for categorical features. Displayed features displayed showed significant association with gene expression subtype (*P*<0.05), except for *CDKN2A* alterations. deltaN expression percentage represents transcript isoform usage between the *TP63* isoforms, deltaN and tap63, as determined by RNA-sequencing. Chromosomal instability (CIN) is defined by the mean of the absolute values of chromosome arm copy numbers from the GISTIC<sup>23,24</sup> output. Absolute values are used so that amplification and deletion alterations are counted equally. Hypermethylation scores and iCluster assignments are described in Supplementary Figure S6.1 and S7.A1, respectively. CIN, methylation, gene expression, and deltaN values were standardized for display using z-score transformation.



**Figure 4. Multi-faceted characterization of mechanisms of** *CDKN2A* **loss a,** Schematic view of the exon structure of *CDKN2A* demonstrating the types of alterations identified in the study. The locations of point mutation are denoted by black and green circles. **b,** *CDKN2A* expression (y-axis) versus *CDKN2A* copy number (x-axis). Samples are represented by circles and colored-coded by specific type of *CDKN2A* alteration. **c,** Diagram of the *KIAA1797-CDKN2A* fusion identified by whole genome sequencing. **d,** *CDKN2A* 

alterations and expression levels (binary) in each sample.

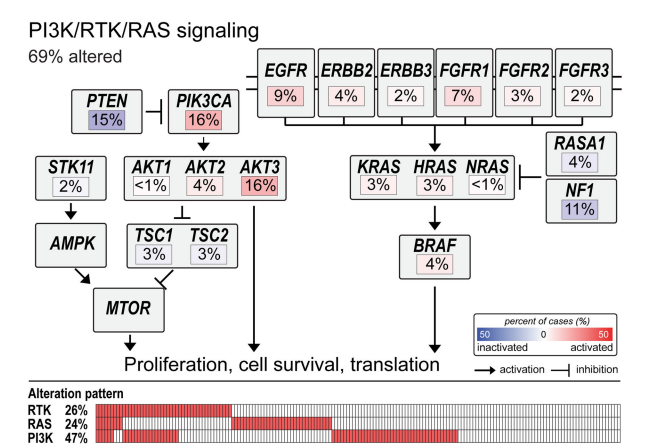


Figure 5. Alterations in targetable oncogenic pathways in lung SqCCs

Pathway diagram showing the percentage of samples with alterations in the PI3K/RTK/RAS pathways. Alterations are defined by somatic mutations, homozygous deletions, high-level, focal amplifications, and, in some cases, by significant up- or down-regulation of gene expression (*AKT3*, *FGFR1*, *PTEN*).