JOURNAL OF CLINICAL ONCOLOGY

Integrative and Comparative Genomic Analysis of Lung Squamous Cell Carcinomas in East Asian Patients

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Published online ahead of print at www.jco.org on December 9, 2013.

Supported by Grant No. A111218-GM08 from the National Project for Personalized Genomic Medicine, Ministry for Health & Welfare, Republic of Korea; by Korea Biobank Network (provided tissue samples; distribution 2012-0006); by Grant No. NCI K08 CA163677 (P.S.H.); and by the Stephen and Alice Cutler Foundation.

Y.K., P.S.H., and J.K. contributed equally to this study. G.G., M.M., and K.P. are principal investigators who contributed equally to this study.

Authors' disclosures of potential conflicts of interest and author contributions are found at the end of this article.

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0732-183X/14/3202w-121w/\$20.00

DOI: 10.1200/JCO.2013.50.8556

Purpose

Lung squamous cell carcinoma (SCC) is the second most prevalent type of lung cancer. Currently, no targeted therapeutics are approved for treatment of this cancer, largely because of a lack of systematic understanding of the molecular pathogenesis of the disease. To identify therapeutic targets and perform comparative analyses of lung SCC, we probed somatic genome alterations of lung SCC by using samples from Korean patients.

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Patients and Methods

We performed whole-exome sequencing of DNA from 104 lung SCC samples from Korean patients and matched normal DNA. In addition, copy-number analysis and transcriptome analysis were conducted for a subset of these samples. Clinical association with cancer-specific somatic alterations was investigated.

Results

This cancer cohort is characterized by a high mutational burden with an average of 261 somatic exonic mutations per tumor and a mutational spectrum showing a signature of exposure to cigarette smoke. Seven genes demonstrated statistical enrichment for mutation: *TP53, RB1, PTEN, NFE2L2, KEAP1, MLL2,* and *PIK3CA*). Comparative analysis between Korean and North American lung SCC samples demonstrated a similar spectrum of alterations in these two populations in contrast to the differences seen in lung adenocarcinoma. We also uncovered recurrent occurrence of therapeutically actionable *FGFR3-TACC3* fusion in lung SCC.

Conclusion

These findings provide new steps toward the identification of genomic target candidates for precision medicine in lung SCC, a disease with significant unmet medical needs.

J Clin Oncol 32:121-128. © 2013 by American Society of Clinical Oncology

INTRODUCTION

Lung cancer is the leading cause of cancer-related death, resulting in an estimated 1.4 million deaths per year worldwide.¹ Lung squamous cell carcinoma (SCC) accounts for approximately 30% of overall lung cancer cases. This type of tumor typically occurs in large airways, and the etiologic role of smoking in lung SCCs is well established.²

In lung adenocarcinoma, it has been observed that patients harboring mutations in the epidermal growth factor receptor (*EGFR*) gene typically demonstrate dramatic responses to EGFR tyrosine kinase inhibitors. This provided the first demonstration that oncogenic alterations could serve as therapeutic biomarkers in lung cancer.^{3,4} Subsequent large-scale clinical trials based on genotype-guided patient stratification have demonstrated a superior clinical benefit of molecularly targeted therapies compared with standard chemotherapy in patients with lung adenocarcinoma who have *EGFR* mutations and *ALK* fusions.⁵⁻⁷ Other molecular inhibitors targeting *BRAF*, *ERBB2*, *RET*, and *ROS* alterations are currently being pursued in clinical trials with promising initial results.

Previous studies have demonstrated recurrent mutation of several genes in SCC, most notably *TP53, NFE2L2,* and *CDKN2A*⁸⁻¹¹ as well as copy-number alterations of *SOX2, CCND1, FGFR1,* and *CDKN2A*¹²⁻¹⁴ in the pathogenesis of lung SCC. Recently, a large-scale comprehensive analysis of a large lung SCC cohort was reported by The Cancer

Genome Atlas (TCGA) Network that identified statistically recurrent somatic mutations in 18 genes that included *TP53*, *NFE2L2*, *KEAP1*, *PIK3CA*, *CDKN2A*, *MLL2*, *HLA-A*, and others.¹⁵ However, in contrast to lung adenocarcinoma, in which molecularly targeted therapies are in routine clinical use to treat subsets of individuals with alterations in genes such as *EGFR* and *ALK*,³⁻⁷ there are no approved targeted therapies for lung SCC. Moreover, subsets of patients with lung cancer with no apparent therapeutic biomarkers, such as EGFR and ALK, generally have worse prognosis, prompting the necessity of identifying driver oncogenic events for this subset of patients.^{1,16}

In lung adenocarcinoma, the prevalence and distribution of specific oncogenic mutations vary widely among demographic groups and are especially pronounced when comparing individuals of mostly white and East Asian descent. For example, East Asians display a higher rate of *EGFR* mutations and a lower rate of *KRAS* mutations.^{17,18} These demographic differences have had a substantial impact on the management of lung adenocarcinoma at regional centers and on the design of clinical trials.

In lung SCC, no large studies have yet been completed to determine the spectrum of genomic alterations in Asians. The prevalence of smoking among Koreans is still high (> 40% among men), and SCC accounts for 25% to 30% of lung cancer cases.¹⁹ To identify therapeutic targets in lung SCC, we probed somatic alterations in tumor samples from Korean patients and compared the results with data from the completely nonoverlapping TCGA cohort,¹⁵ which consisted primarily of patients from academic centers in the United States. Here, we present the multidimensional analysis of genomic alterations in more than 100 Korean patients with lung SCC by using whole-exome and transcriptome sequencing and single nucleotide polymorphism array analysis.

PATIENTS AND METHODS

Study Population

The study included an institutional review board–approved retrospective analysis of 104 Korean patients with lung SCC. Information on sex, age, tumor stage, smoking record, and overall survival was extracted from the clinical database for this cohort (Data Supplement). None of these patients had previously been subject to genomic characterization, and no patients from these hospitals were part of the TCGA study of lung SCC.

Sample Preparation, Sequencing Data Analysis, and Statistical Methods

Analysis of sequencing data was performed as previously described.^{15,20} Detailed methods for sample preparation and sequencing analysis can be found in the Data Supplement (for statistical analysis, each statistical method used is described in the text where the data are presented). Enrichment analysis of specific somatic mutations and copy-number alterations in somatic copy-number alteration (SCNA) clusters was performed by using Fisher's exact test. Exclusivity and co-occurrence of mutations were examined by using one-tailed Fisher's exact tests. For multiple test adjustment, false discovery rate was controlled by calculating *q* values.²¹ Other statistical methods are described in the main text and the Data Supplement.

RESULTS

Patient Cohort Description

To provide comprehensive understanding of genetic abnormalities occurring in lung SCC, we used massively parallel paired-end

Table 1. Clinical Data Summary				
	Korean (n = 104)		TCGA ¹⁵ (n = 178)	
Variable	No.	%	No.	%
Age at surgery, years Median Range	65 43-86		68.0 40.0-85.0	
Sex Male Female	100 4	96.2 3.8	131 47	73.6 26.4
Smoking status Never-smoker Former smoker, pack-years ≤ 20 > 20	5 52 7 45	4.8 50 6.7 43.3	7 137	3.9 77
Current smoker, pack-years ≤ 20 > 20 N/A	47 4 43 0	45.2 3.8 41.3 0	28	15.7 3.4
Median follow-up, months	29		15.8	
Tumor stage I II III IV N/A	43 41 18 2 0	41.3 39.4 17.3 1.9 0	97 38 38 3 2	54.5 21.3 21.3 1.7 1.1
T stage T1 T2 T3 T4 N/A	24 63 12 5 0	23.1 60.6 11.5 4.8 0	35 116 14 12 1	19.7 65.2 7.9 6.7 0.6
N stage N0 N1 N2 N3	70 20 14 0	67.3 19.2 13.5 0	117 39 17 5	65.7 21.9 9.6 2.8

Abbreviations: N/A, not applicable; TCGA, The Cancer Genome Atlas.

sequencing technology²² to perform whole-exome solution-based hybrid capture sequencing of 104 matched tumor/normal pairs. These samples were from individuals who underwent resection of a lung SCC from 2005 to 2012. The median age of patients in the study was 65 years with a median of 29.0 months of follow-up (Korean columns in Table 1). The cohort included five self-reported never-smokers, 52 former smokers, and 47 current smokers; most patients had extensive smoking histories. The cohort included 43 stage I, 41 stage II, 18 stage III, and two stage IV lung SCC tumors, all of which were primary resection specimens. Clinical and histopathologic information for the cohort set used in the study is provided in the Data Supplement. In addition to whole-exome sequencing, we also conducted transcriptome sequencing and copy-number analysis for a subset of this patient cohort (Data Supplement).

Detection of Somatic Aberrations

Analysis of whole-exome sequencing data from our cohort identified a total of 29,353 point mutations and 935 insertion or deletion events across 104 tumor/normal pairs in coding regions of the genome. Mean sequencing depth in the target regions was 62.8X. Of the 29,353 point mutations, 21,550 resulted in an amino acid change in the corresponding proteins. Individuals displayed a mean point mutation rate of 8.71 mutations per Mb (range, 1.19 to 36.2 mutations per Mb). Among nonsynonymous substitutions, transitions and transversions at CpG sites were the most commonly observed mutational context with a rate of 16.18 per Mb and 9.53 per Mb, respectively. Mutations at Cp(A/C/T) sites were the second most commonly observed context at 7.68 per Mb (Data Supplement). Hierarchical clustering of tumors based on mutational spectra is shown in the Data Supplement.

We applied the MutSig method to identify genes that demonstrated evidence of statistical selection for mutation in our cohort.^{20,23-25} MutSig is a computational algorithm that takes into account gene length, composition, silent-to-nonsilent mutation ratios, and other factors, and it identified six genes that displayed significant enrichment for mutations as defined by q < 0.1, with robust evidence of expression in lung SCC: TP53, RB1, PTEN, NFE2L2, KEAP1, and MLL2 (Fig 1 and Data Supplement). A second MutSig analysis, in which only genes annotated in the Catalogue of Somatic Mutations in Cancer (COSMIC) database were considered, rediscovered TP53, RB1, and PTEN and additionally nominated PIK3CA as statistically significant (Data Supplement). Additional genes with annotated roles in cancer that did not reach statistical significance but did display recurrent mutations were CHD7 (nine mutations), KRAS (two), NF1 (13), EGFR (two), CDKN2A (three), IBTK (three), TIE1 (five), NOTCH1 (seven), NOTCH3 (nine), FBXW7 (four), FGFR1 (two), FGFR2 (two), KDM6A (five), and APC (four; Fig 1). Of note, one of two patients with an EGFR mutation demonstrated a canonical exon 19 deletion and the other demonstrated D347N. Both KRAS mutations resulted in G12C substitutions.

SCNAs were inferred from sequencing data by read-depth analysis (Data Supplement). We used GISTIC 2.0 analysis^{26,27} to identify statistically recurrent peaks of amplification and deletion and identified 19 regions of focal amplification and 25 regions of deletion with a q < 0.25 (Data Supplement). The most significantly amplified regions were 3q26, an amplicon including *SOX2*; 8p11 (*FGFR1, WHSC1L1*); 7p11 (*EGFR*); 11q13 (*CCDN1*); and 4q12 (*KDR, KIT, PDGFRA*). Deletion peaks tended to be more broad and included 2q37, 4q35 (*CASP3*), 9p21 (*CDKN2A*), and 10q23 (*PTEN*). The copy-number variations were also confirmed and validated by using high-density single nucleotide polymorphism arrays (Data Supplement).

Hierarchical clustering of the cohort based on the SCNAs demonstrated three clusters (Data Supplement): Cluster 1 was defined by a paucity of SCNAs (depicted in red in Fig 1C), Cluster 2 was defined by both broad and focal 3q amplification (depicted in green in Fig 1C), and Cluster 3 was defined by broad 3q amplification and SCNAs in cyclin genes and receptor tyrosine kinase-related genes (depicted in magenta in Fig 1C; Fisher's exact test: CCND1, P < .03; CCND2, P < .001; *ERBB*2, *P* < .04; *FGFR*1, *P* < .02; *EGFR*, *FRS2*, and *PDGFRA*, all P < .001). Cluster 1 demonstrated a lower mutation rate compared with the other clusters (Data Supplement). This cluster also contained some samples with lower calculated tumor purity, which may have contributed to a lower prevalence of genomic alterations. Cluster 2 exhibited near exclusivity with RB1 mutations (Fisher's exact test: P < .001) and enriched in *PIK3CA*-activating mutations (P < .007). Virtually all the patients in the third cluster (magenta in Fig 1C) possessed *TP53* mutations (Fisher's exact test: P < .001; Data Supplement).

Paired-end RNA sequencing was performed on a subset of 26 individuals from the original cohort. We first used these data to validate our mutation calls by calling the mutations in the RNA sequencing data. After controlling for coverage, we validated more than 72% of coding single nucleotide variants and 86% of events in genes depicted in Figure 1 (Data Supplement). Analysis of these data identified a total of 296 gene fusions involving 244 distinct fusion partners (Data Supplement).²⁸ No fusions involving the *ALK*, *RET*, or *ROS1* kinase genes were observed. We observed recurrent fusions involving the *CASP1* gene (four), *IGF2BP2* (two), and *KRT5* (two). We identified single-fusion events involving the *FGFR3* and *GSK3* β kinases in the 26 initial individuals surveyed.

Detailed analysis of the FGFR3 event (in sample J9) demonstrated an in-frame fusion of exon 8 of TACC3 with exon 17 of FGFR3 (Fig 2A), a fusion event recently reported as oncogenic and potentially therapeutically targetable in glioblastoma multiforme, bladder cancer, and other types of cancer, including prostate cancer.²⁹⁻³³ We extended analysis to an additional cohort of 40 patients with lung SCC and identified an additional individual (J1) with an FGFR3-TACC3 fusion by reverse transcriptase polymerase chain reaction (Fig 2B). We conducted genomic DNA polymerase chain reaction and Sanger sequencing and identified the genomic breakpoints and joining points for these two samples containing FGFR3-TACC3 fusion transcripts (Data Supplement). Given these observations, we further probed the TCGA lung SCC RNA sequencing data set15 and identified four additional patients who had tumors containing FGFR3-TACC3 in-frame fusion transcripts of 178 tumor RNA specimens (Data Supplement), further confirming that these fusion events are recurrent in lung SCC. All six patients with fusion transcripts involved the omission of the last exon of FGFR3, which was instead fused in-frame to various exons of TACC3 and accompanied by an increase in FGFR3 transcript level (Data Supplement).

Comparative Mutational Analysis of Lung SCCs in East Asian and Mostly White Patients From the United States

By using the set of genomic alterations in this lung SCC cohort, we investigated the association among the spectrum of somatic mutations with clinical parameters. Most of the major somatic mutations did not have a statistical association with overall survival (Data Supplement). We did observe that some mutations or SCNAs were significantly associated with poor survival (Data Supplement). The biologic significance of alterations in genes with apparent prognostic value, such as *MLL2* mutations, requires further experimental analysis and validation in additional independent cohorts.

Given that no systematic comparison has yet been reported for lung SCC among different ethnicities, we compared the spectra of mutational and SCNAs of this cohort to a similarly sized cohort recently reported by the TCGA,¹⁵ consisting mostly of white patients from the United States (Table 1). Both cohorts displayed a similarly high rate of mutations of *TP53* (73% Korean, 81% TCGA; Fisher's exact P > .16), mutations in the *NFE2L2/KEAP1* genes (33% and 31%; P > .85), and those in *PIK3CA/PTEN* (24% and 24%) and *NOTCH1* (7% and 8%; P > .9). Both cohorts also demonstrated enrichment for CpG transition and transversion mutations. SCNAs involving *SOX2*, *FGFR1*, *KDR/PDGFRA/KIT*, and *PTEN* were also observed at comparable frequencies. One notable difference among

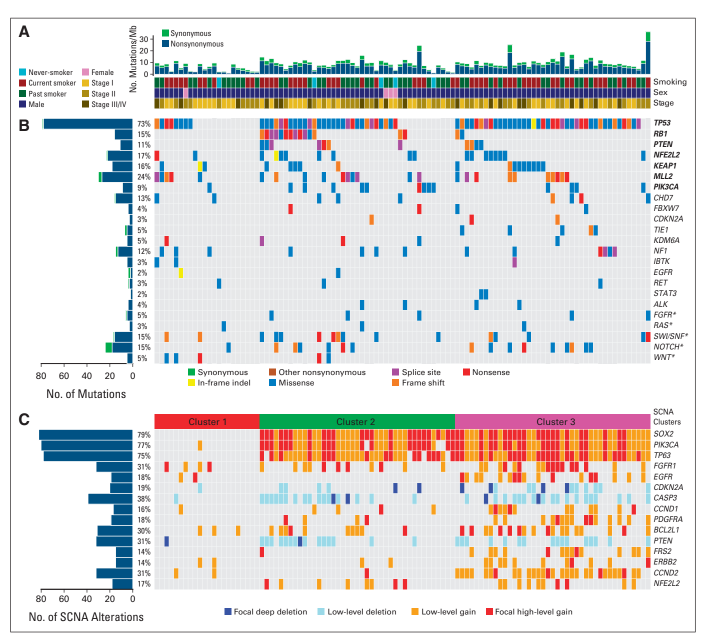


Fig 1. Clinicopathologic characteristics and genetic aberrations across 104 lung squamous cell carcinomas. (A) Rate of synonymous and nonsynonymous mutations, expressed in mutations per megabase (Mb) of covered target sequence in association with clinical parameters for the samples. In the bottom panel, the first row indicates smoking status, the second row sex, and the third row cancer stage. (B) Heatmap representation of individual mutations present in 104 lung squamous cell carcinoma samples in association with information from (A). It displays the mutational types in a given sample and in a given gene in a two-dimensional matrix. (Left) Percentages of mutations, representing at least one mutational event in each gene. (Right) List of recurrently mutated genes. *FGFR, RAS, SWI/SNF, NOTCH*, and *WNT* mini-gene sets are shown as single tracks because there was complete exclusivity of mutations within these gene families. *FGFR** indicates conglomerated representation of *FGFR1 + FGFR3, RAS*: HRAS + KRAS, SWI/SNF*: ARID1A + ARID1B + PBRM1, NOTCH*: NOTCH1 + NOTCH3, WNT*: APC + CTNNB1.* (C) Somatic copy-number alterations (SCNAs) of samples in association with (A) and (B). Significant SCNAs are shown, highlighting the samples with threshold copy-number changes specified in the lower bottom panel. Indel, insertions and deletions.

the data sets was somatic alterations in cell cycle regulatory genes, including *CDKN2A* and *RB1*. In the TCGA cohort, *CDKN2A* mutations were observed in 15% of patients and homozygous deletions in 33%. In contrast, *CDKN2A* mutations and copy-number alterations were infrequent in the Korean cohort (4% and 5%, respectively; P < .007 and P < .001, respectively); however, mutations of *RB1* were more common in the Korean cohort (15%) than in the TCGA cohort (4%; Fisher's exact P < .005).

Both our data and the TCGA data set¹⁵ confirmed previous reports suggesting that the two most common oncogenic mutations in lung adenocarcinomas, *KRAS* and *EGFR*,³⁴ are rare in lung SCCs. We identified only one case with an activating *EGFR* mutation in this cohort, an exon 19 deletion, and no cases of L861Q, which was observed in the TCGA cohort.¹⁵ We noted two cases of *KRAS* G12C but no codon 12 or 13 *RAS* events were reported by TCGA (Data Supplement). Alterations in other *RAS* family members such as *NRAS* and

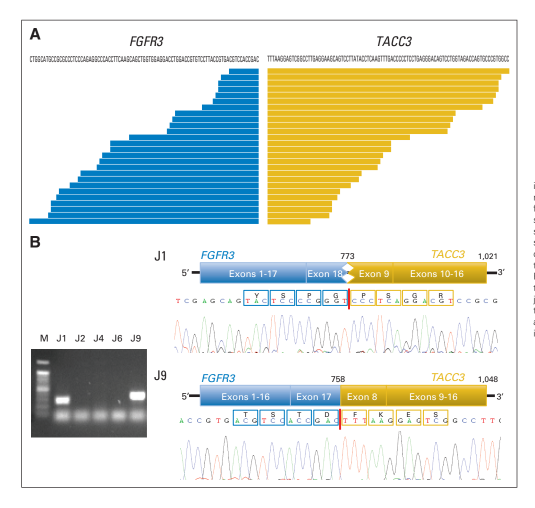


Fig 2. Recurrent *FGFR3-TACC3* fusion identified in lung squamous cell carcinoma. (A) *FGFR3-TACC3* fusion identified from RNA sequencing; 26 split-reads that span the *FGFR3-TACC3* junction are shown. (B) Left: *FGFR3-TACC3* fusion-specific polymerase chain reaction from complementary DNA derived from patients with lung squamous cell carcinoma. Right: sequencing chromatogram from two patient samples spanning the fusion junction. Shown in box are the translational reading frames and designation of amino acids. M, DNA marker. J1, patient identification number.

HRAS were virtually absent in this cohort, although putative loss-offunction events in negative regulators of *RAS* such as *RASA1* and *NF1* were observed at appreciable frequencies (7% and 13%, respectively; summarized in the Data Supplement).

Closer examination of mutational profiles revealed core signaling pathways implicated in discrete functional categories. As in the TCGA cohort analysis, a transcription factor pathway regulating redox stress comprising NFE2L2, KEAP1, and CUL3 aberrations was altered in 40% of the patients (Fig 3). Another pathway implicated was deregulated chromatin remodeling with alterations in chromatin remodeling enzymes, including histone demethylases and SWI-SNF complexes identified in more than 49% of patients (Fig 3). Some members in this group of core pathways demonstrated mutually exclusive mutational patterns (Fig 3). An exception to this exclusivity relationship was the strict co-occurrence of MLL2 and KDM6A mutations (Fisher's exact P < .001). It is of note that members of these two core pathways were also likely to interact functionally, given the observation that mutations in NFE2L2 and ARID1A or ARID1B co-occur (P < .001; Data Supplement). Squamous cell differentiation pathways involving inactivating mutations of NOTCH genes were also frequently altered^{15,20} (Fig 3). FGFR kinase amplification and mutations were frequent and were identified in 18 individuals, and PI3K was the most commonly altered serine/threonine kinase family with mutational events in 17 individuals (Fig 3).

DISCUSSION

This study has identified a potentially targetable fusion gene in lung SCCs, FGFR3-TACC3, which defines a new specific subset for potential targeted therapy in lung SCC and was recently reported by others.^{32,33} Previously, FGFR small-molecule inhibitors were shown to possess therapeutic activity in lung SCC cells with FGFR family kinase amplification and mutation.^{14,35} Clinical activity and safety of BGJ398 in FGFR1-amplified lung SCC and AZD4547 in various types of solid tumors have recently been demonstrated.^{36,37} The FGFR antagonists are also under clinical investigation in lung SCCs with mutations or amplification of FGFR1, FGFR2, or FGFR3 as more diversified inclusion criteria (clinical trials NCT01761747 and NCT01795768). Preclinical model studies of FGFR3-TACC3 in glioblastoma and bladder cancer demonstrated encouraging utility of FGFR inhibitors in the treatment of cancer cells harboring the fusion gene.²⁹⁻³¹ Thus, it is likely that clinical trials could be expanded to test the efficacy of these agents in patients with FGFR3 fusions in lung SCC with concurrent development of companion diagnostics. Despite apparently modest demographic influences in lung SCC, this study also points out the value of analyzing different cohorts, since the TCGA study initially missed the identification of the FGFR3 fusion events.15

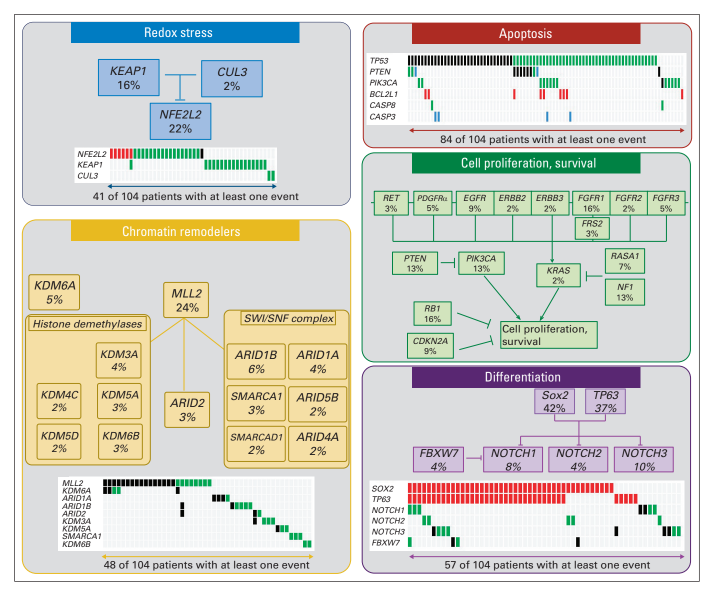


Fig 3. Diagram of integrated lung squamous cell carcinoma pathways. Proposed diagram integrating five different core pathways of lung squamous cell carcinoma. Green, substitution mutations; black, truncating mutations; red, amplifications; blue, deletions.

In contrast to lung adenocarcinomas in which the frequency of genomic alterations in oncogenes differs among ethnic groups,^{34,38} in lung SCC we found that the mutational landscape of two different population groups with similar sized samples had overall congruent mutational frequencies in major oncogenic drivers and tumor suppressors. To the best of our knowledge, this is the first and largest comparison of the integrative genomic analysis of a given cancer type with more than 100 individuals between populations of Asians and mostly white patients from the United States. Although the frequency of mutations in RB1 and CDKN2A genes were significantly different between the two cohorts, frequencies of other major genes, including TP53, NFE2L2, KEAP1, PIK3CA, PTEN, MLL2, and NOTCH1, were similar to each other. Thus, in lung SCC, in which most of the patients had been chronically exposed to heavy doses of tobacco, the subtle genetic and inherent biologic influences seem to be largely overridden by heavy mutational burdens from exogenous carcinogens. It has been shown that several cancer types, including melanoma, display highly increased mutational burden with exogeneous carcinogens linked to their tumorigenesis. Systematic comparative genomic profiling of this and other cancer types with high mutational rates in patients in different ethnic groups will help address the generalizability of the findings.

Although most SCCs occur in smokers, a small subset of neversmoking patients (five of 104 for this study; seven of 178 for TCGA) still succumb to the disease. We did not capture information regarding exposure to second-hand smoke or environmental toxins such as asbestos in this study, and smoking status was self-reported. We believe that larger-scale genomic studies specifically targeting these subsets will help to better elucidate underlying biologic mechanisms associated with the never-smoker lung SCC subgroup. In this sense, it is of note that ongoing large-scale cancer consortium studies, including that of TCGA, currently aim to delineate genomic aberrations in more than 500 patients for each subtype of cancer in the coming years. The sample size used in this study (n = 104; n = 178 for TCGA) and the lower degree of sequencing coverage compared with that reported in TCGA limit the ability to perform comparative analyses of less common clinical features or genomic alterations with sufficient statistical power. Given the complexities of the cancer genomic landscape, it is likely that expansion of patient samples in the future will help to further clarify subtle genetic differences in lung SCC contributed by ethnicity. It should also be noted that demographic heterogeneity within the TCGA cohort was not well annotated, which prevented the possible exclusion of patients of Asian descent in this cohort for our comparative analyses.

In summary, this study has comparatively described the recurrent somatic mutations in lung SCCs in a large cohort of East Asians. It revealed the commonality in the pathogenesis of lung SCC across ethnicities and identified several targetable oncogenic mutations in a disease with large unmet medical needs.

AUTHORS' DISCLOSURES OF POTENTIAL CONFLICTS OF INTEREST

Although all authors completed the disclosure declaration, the following author(s) and/or an author's immediate family member(s) indicated a financial or other interest that is relevant to the subject matter under consideration in this article. Certain relationships marked with a "U" are those for which no compensation was received; those relationships marked with a "C" were compensated. For a detailed description of the disclosure categories, or for more information about ASCO's conflict of interest policy, please refer to the Author Disclosure Declaration and the Disclosures of Potential Conflicts of Interest section in Information for Contributors.

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Employment or Leadership Position: None Consultant or Advisory Role: Matthew D. Wilkerson, GeneCentric Diagnostics (C); Matthew Meyerson, Bayer Pharmaceuticals (U), Foundation Medicine (C) Stock Ownership: Matthew Meyerson, Foundation Medicine Honoraria: None Research Funding: None Expert Testimony: None Patents: Michael S. Lawrence, MutSigCV; Matthew Meyerson, patent US20060147959 Other Remuneration: Michael S. Lawrence, MutSigCV

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Final approval of manuscript: All authors

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Acknowledgment

We thank William Hahn, MD, PhD (Dana-Farber Cancer Institute) for his generous advice and input; Macrogen for sequencing efforts; Theragen Bio for quality control efforts; C. Maeng, H. Jung, S. Park, J. Cha, W. Shin, and E. Lee (Samsung Medical Center) for their valuable input into the project; Y.S. Lee for her excellent graphical help; and PGM21 and its director, H. Kim, MD, for their continued support of the project.