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with Massively Parallel Sequencing*

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Mapping the Hallmarks of Lung Adenocarcinoma with Massively Parallel Sequencing

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SUMMARY

Lung adenocarcinoma, the most common subtype of non-small cell lung cancer, is responsible for more than 500,000 deaths per year worldwide. Here, we report exome and genome sequences of 183 lung adenocarcinoma tumor/normal DNA pairs. These analyses revealed a mean exonic somatic mutation rate of 12.0 events/megabase and identified the majority of genes previously reported as significantly mutated in lung adenocarcinoma. In addition, we identified statistically recurrent somatic mutations in the splicing factor gene *U2AF1* and truncating mutations affecting *RBM10* and *ARID1A*. Analysis of nucleotide context-specific mutation signatures grouped the sample set into distinct clusters that correlated with smoking history

and alterations of reported lung adenocarcinoma genes. Whole-genome sequence analysis revealed frequent structural rearrangements, including in-frame exonic alterations within *EGFR* and *SIK2* kinases. The candidate genes identified in this study are attractive targets for biological characterization and therapeutic targeting of lung adenocarcinoma.

INTRODUCTION

Lung cancer is a leading cause of death worldwide, resulting in more than 1.3 million deaths per year, of which more than 40% are lung adenocarcinomas (World Health Organization, 2012; Travis, 2002). Most often, tumors are discovered as locally advanced or metastatic disease, and despite improvements in molecular diagnosis and targeted therapies, the average 5 year

survival rate for lung adenocarcinoma is ~15% (Minna and Schiller, 2008).

Molecular genotyping is now routinely used to guide clinical care of lung adenocarcinoma patients, largely due to clinical trials that demonstrated superior efficacy of targeted kinase inhibitors as compared to standard chemotherapy for patients with *EGFR* mutations or *ALK* fusions (Kwak et al., 2010; Pao and Chmielecki, 2010). In addition to *EGFR* and *ALK* alterations found in ~15% of U.S. cases, lung adenocarcinomas frequently harbor activating mutations in *KRAS*, *BRAF*, *ERBB2*, and *PIK3CA* or translocations in *RET* and *ROS1* (Pao and Hutchinson, 2012), all of which are being pursued as targets in ongoing clinical trials (<http://clinicaltrials.gov>). Lung adenocarcinomas also often harbor loss-of-function mutations and deletions in tumor suppressor genes *TP53*, *STK11*, *RB1*, *NF1*, *CDKN2A*, *SMARCA4*, and *KEAP1* (Ding et al., 2008; Kan et al., 2010; Sanchez-Cespedes et al., 2002). Unfortunately, such alterations are difficult to exploit therapeutically. Therefore, knowledge of additional genes altered in lung adenocarcinoma is needed to further guide diagnosis and treatment.

Previous efforts in lung adenocarcinoma genome characterization include array-based profiling of copy number changes (Tanaka et al., 2007; Weir et al., 2007), targeted sequencing of candidate protein-coding genes (Ding et al., 2008; Kan et al., 2010), and whole-genome sequencing of a single tumor/normal pair (Ju et al., 2012; Lee et al., 2010). These studies identified somatic focal amplifications of *NKX2-1*, substitutions and copy number alterations in known oncogenes and tumor suppressor genes, and recurrent in-frame fusions of *KIF5B* and *RET*. These studies have also nominated several putative cancer genes with somatic mutations (*EPHA* family, *NTRK* family, *TLR4*, *LPHN3*, *GRM1*, and *GLI*), but the functional consequence of many alterations is unknown. A recent study describing whole-exome sequencing of 16 lung adenocarcinomas (Liu et al., 2012) enumerated several mutated genes but did not identify genes undergoing positive selection for mutation in the studied tumors.

In this study, we used next-generation sequencing to sequence the exomes and/or genomes of DNA from 183 lung adenocarcinomas and matched normal adjacent tissue pairs. In addition to verifying genes with frequent somatic alteration in previous studies of lung adenocarcinoma, we identified novel mutated genes with statistical evidence of selection and that likely contribute to pathogenesis. Together, these data represent a significant advance toward a comprehensive annotation of somatic alterations in lung adenocarcinoma.

RESULTS

Patient Cohort Description

We sequenced DNA from 183 lung adenocarcinomas and matched normal tissues by using paired-end massively parallel sequencing technology (Bentley et al., 2008). The cohort included 27 never-smokers, 17 light smokers (defined by less than ten pack years of tobacco use), 118 heavy smokers (more than ten pack years), and 21 patients of unknown smoking status (Table 1). The cohort included 90 stage I, 36 stage II, 22 stage III, and 10 stage IV lung adenocarcinoma cases, as well as 25 patients with unknown stage. All tumors were chemotherapy-

Table 1. Summary of Clinical Features

Age at Surgery (Median; Range)	66 (36–87)
Gender	
male	95
female	88
Smoking Status (AJCC 7th Edition)	
never-smoker	27
smoker >10 years	118
smoker ≤ 10 years	17
n/a	21
pack years (median; range)	30 (0–128)
Survival	
follow-up available	135
follow-up unavailable	48
PFS in months (median; range)	9 (0–63)
Tumor Stage	
I	90
II	36
III	22
IV	10
n/a	25
Distribution of selected clinical variables from 183 lung adenocarcinoma cases.	

naive, primary resection specimens except for one case with whole-genome sequence data (LU-A08-43) that was a postchemotherapy metastatic tumor from a never-smoker. Sample acquisition details are provided in [Extended Experimental Procedures](#). Additional clinical descriptors of the cohort are provided in [Table 1](#). Comprehensive clinical and histopathological annotations, sequence characteristics, and major variants for each patient in the study are provided in [Table S1](#) (available online).

Mutation Detection and Validation

We examined 183 lung adenocarcinoma tumor/normal pairs with a combination of whole-exome sequencing (WES) or whole-genome sequencing (WGS): 159 WES, 23 WES and WGS, and 1 WGS only. Exomes were sequenced to a median fold coverage of 92 (range: 51–201) on 36.6 Mb of target sequence (Fisher et al., 2011). Genomes were sequenced to a median coverage of 69 (range: 25–103) in the tumor and 36 (range: 28–55) in the normal, with the higher tumor coverage to adjust for stromal contamination. Complementary SNP array analysis of 183 pairs was used to detect genome-wide somatic copy number alterations. See [Table 2](#) and [Extended Experimental Procedures](#) for more details.

We identified somatic substitutions and small insertions and deletions (indels) through statistical comparison of paired tumor/normal sequence data by using algorithms calibrated for stromally contaminated cancer tissues (Banerji et al., 2012; Stransky et al., 2011) (www.broadinstitute.org/cancer/cga; [Extended Experimental Procedures](#)). Exonic regions of the 183 cases contained 77,736 somatic variants corresponding to

Table 2. Whole-Genome and Whole-Exome Sequencing Statistics

Statistic	Whole-Exome	
	Capture	Whole Genome
Tumor/normal pairs sequenced	159	24
Total tumor Gb sequenced	1,031.6	4,946.0
Median fold tumor target coverage (range)	91 (51–201)	69 (25–103)
Median normal fold target coverage (range)	92 (62–141)	36 (28–55)
Median somatic mutation rate per Mb in target territory (range)	6.8 (0.3–94.7)	13.3 (4.5–55.3)
Median number of coding mutations per patient (range)	216 (1–3,512)	323 (63–2,279)
Median number of nonsynonymous mutations per patient (range)	167 (1–2,721)	248 (53–1,770)
Median number of transcribed noncoding mutations per patient (range)	187 (13–2,559)	18,314 (4,632–100,707)
Total number of structural rearrangements	n/a	2,349
Total number of frame-preserving genic rearrangements	n/a	71
Total number of frame-abolishing genic rearrangements		235
Median number of genes powered at 20% exonic territory (range)	15,647 (15,046–16,019)	16,905 (10,136–16,952)
Median number of genes powered at 50% exonic territory (range)	6,788 (6,078–7,402)	8,771 (2,634–8,863)

Selected sequencing statistics for 183 WES and WGS cases. “Tumor Target Territory” refers to the exonic territory targeted by the exome capture bait set reported by (Fisher et al., 2011) and used in this study. The “Whole-Exome Capture” column does not include data on 23 cases analyzed by both WES and WGS.

a median of 8.1 mutations/Mb and a mean of 11.9 mutations/Mb (range: 0.04–117.4). These comprised 43,813 missense, 14,801 silent, 3,504 nonsense, 1,460 splice-site, 2,310 deletions, 839 insertions, and 11,009 other mutations (predominantly residing in 5' and 3' untranslated regions [UTRs]). Of the 3,149 indels, 182 were in-frame, 1,785 were predicted to cause a frame shift, 68 occurred at a splice site, and 1,114 were otherwise classified.

Mutation calls were validated by cross-comparison of coding mutations detected by WES and WGS from 24 cases with both data types. We validated 84% of 380 indels and 97% of 9,354 substitutions identified by WGS at sufficiently powered sites in the corresponding WES tumor sample. In the converse analysis, we validated 86% of 338 indels and 98% of 8,912 substitutions from WES at sufficiently powered sites in the corresponding

WGS tumor sample (Figure S1, Table S2A, and Extended Experimental Procedures). To validate mutations from cases with only WES data, we randomly selected 69 candidate mutations for ultradeep (>1,000-fold) targeted resequencing. Somatic status was confirmed for 30 of 33 (91%) indel events and 33 of 36 (92%) substitution events (Table S2B). These validation rates generally meet or exceed those reported in similar sequencing studies (Banerji et al., 2012; Berger et al., 2012; Gerlinger et al., 2012; Nikolaev et al., 2012; Stransky et al., 2011; Cancer Genome Atlas Research Network, 2011; Totoki et al., 2011; Zang et al., 2012).

Somatic Genetic Signatures of Mutagen Exposure in Lung Adenocarcinoma

Consistent with previous studies (Ding et al., 2008; Kan et al., 2010; Zang et al., 2012), we observed significantly higher exonic mutation rates in tumors from smokers (median: 9.8/Mb; mean: 12.9/Mb; range: 0.04–117.4/Mb) compared to never-smokers (median: 1.7/Mb; mean: 2.9/Mb; range: 0.07–22.1/Mb; $p = 3.0 \times 10^{-9}$, Wilcoxon rank sum test). Lung adenocarcinoma mutation rates in our cohort exceeded those reported for other epithelial tumor types, except melanoma and squamous cell lung cancer (Hodis et al., 2012; Nikolaev et al., 2012; Cancer Genome Atlas Research Network, 2012; Wei et al., 2011).

To characterize the mutation spectrum of lung adenocarcinoma, we analyzed somatic substitutions and covered bases within their trinucleotide sequence context (Figure 1A). The most frequent mutation signatures were C→T transitions in the setting of CpG dinucleotides (CpG→T) and C→A transversions. The least frequent mutation type was A→C. Unbiased hierarchical clustering of context-specific mutation rates across 182 WES cases yielded five mutation spectrum clusters. These clusters represented grades of increasing mutational complexity: cluster 1 was enriched for CpG→T mutations and was marked by an overall low mutation rate; cluster 2 was characterized by CpG→T transitions and CpG→A transversions; cluster 3 showed additional C→A transversions outside of the CpG context; cluster 4 showed additional C→T transitions outside of the CpG context and TpC transversions that mutated to either a T or a G; and cluster 5 comprised hypermutated tumors containing a broad mutational spectrum that included rare mutation signatures, such as A→T transversions. Mutation spectrum clusters in tumors correlated with clinical features of patients. Cluster 1 was significantly enriched in never- and light smokers ($p = 1.9 \times 10^{-9}$, Fisher's exact test), whereas cluster 4 was significantly enriched in patients with advanced (IIIB or IV) stage ($p = 0.0063$, Fisher's exact test).

Differentiation of smokers and never-smokers was evident from comparison of mutation counts from the most frequent mutational signatures, CpG→T and C→A (Figure 1B). These results were consistent with previous reports of signatures of DNA damage by tobacco (Hainaut and Pfeifer, 2001). Applying thresholds to a log-adjusted ratio of CpG→T and C→A mutations (see Experimental Procedures), we imputed smoking status for 21 patients who lacked reported smoking history and accurately recapitulated reported smoking status for more than 75% of the remaining cases (Figure 1B). Exonic and intronic mutation rates, context-specific mutation counts, imputed

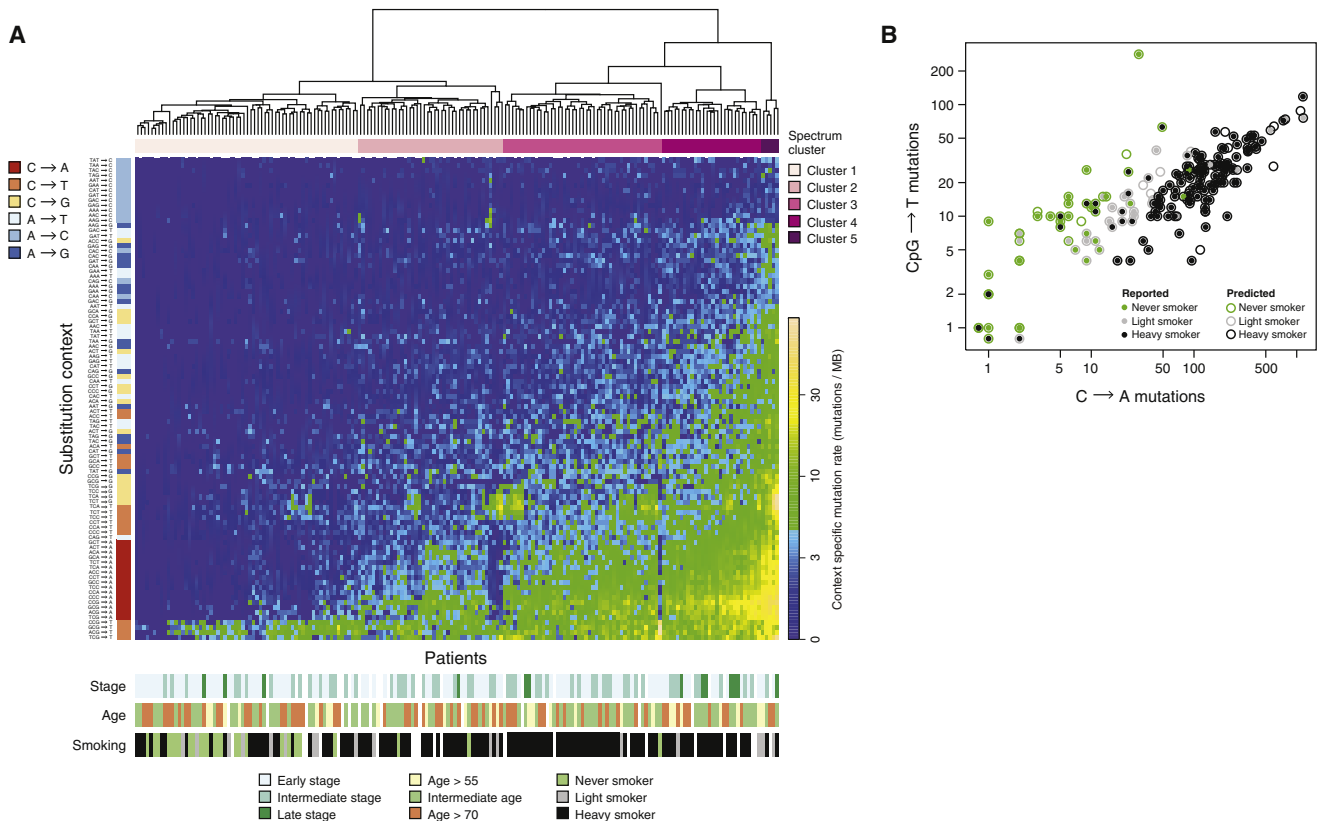


Figure 1. Mutation Spectrum Analysis of 183 Lung Adenocarcinomas

(A) Hierarchical clustering of 183 lung adenocarcinomas according to their nucleotide context-specific exonic mutation rates. Each column represents a case, and each row represents one of 96 strand-collapsed trinucleotide context mutation signatures. Top bar, patient-cluster membership; left bar, simplified single-nucleotide context mutational signature; bottom bars, reported tumor stage, age, and smoking status for each patient; right gradient, mutation rate scale.

(B) Stratification of reported versus imputed smoking status by the log transform of the adjusted ratio of C \rightarrow A transversion rates and CpG \rightarrow T transition rates. The color of each inner solid point represents the reported smoking status for that particular patient. The color of each outer circle indicates that patient's imputed smoking status as predicted by the classifier. Additional analytic details are provided in the [Extended Experimental Procedures](#).

See also [Figure S1](#) and [Tables S2](#) and [S3](#).

smoking status, and mutation spectrum cluster assignments for each patient are provided in [Table S1](#).

Calibration of a Statistical Approach to the Analysis of High Mutation Rate Tumors

The high mutation rates in lung adenocarcinoma and other tumors (Hodis et al., 2012; [Cancer Genome Atlas Research Network, 2012](#)) present a challenge for unbiased discovery of mutated genes undergoing positive somatic selection. More than 13,000 of 18,616 genes with adequate sequence coverage had nonsynonymous somatic mutations in at least one tumor, and more than 3,000 were mutated in at least five patients. These genes included those with very large genomic footprints (e.g., *TTN*), genes with low basal expression in lung adenocarcinomas (e.g., *CSMD3*), and genes accumulating high numbers of silent substitutions (e.g., *LRP1B*).

Application of a standard binomial background mutation model assuming a constant mutation rate in each patient and nucleotide context stratum (Berger et al., 2011) yielded profound test statistic inflation ([Figure S2A](#)) and identified more than 1,300

significantly mutated genes. Genes with significant p values in this analysis had low basal expression in lung adenocarcinoma cell lines (Barretina et al., 2012) ([Figure S2B](#)), harbored high fractions of synonymous mutations, and were enriched in gene classes previously unassociated with cancer (e.g., olfactory receptors and solute transporters). Recalibration of this model by limiting to genes with evidence of expression improved, but did not completely correct, this statistical inflation ([Figure S2C](#)). These results suggested a high degree of variation in neutral somatic mutation rates among genes, including expression-dependent variation. This observation is consistent with reports of regional mutation rates correlated with density of H3K9 chromatin marks across cancers (Schuster-Böckler and Lehner, 2012) and with gene expression in multiple myeloma (Chapman et al., 2011).

To more adequately model variation of neutral somatic mutation rates among genes, we applied the InVEx algorithm (Hodis et al., 2012) to exploit the abundant noncoding mutations detected by both WES and WGS. InVEx permutes coding, untranslated, and intronic mutations within covered territories of each

gene, patient, and nucleotide context to generate within-gene null distributions of “functional impact” across a sample set (see [Experimental Procedures](#)).

Our primary InVEx analysis employed a PolyPhen-2 (PPH2)-based metric ([Adzhubei et al., 2010](#)) to assess the functional impact of observed and permuted mutations. Applying this analysis to 12,907 mutated genes with at least one PPH2-scored event yielded a well-distributed test statistic with minimal inflation ([Figure S2D](#)) and without gene expression bias in lung adenocarcinoma cell lines ([Figure S2B](#)). To increase specificity and power, we restricted our analysis to 7,260 genes demonstrating expression (median Robust Multiarray Average [RMA] value ≥ 5) in a panel of 40 lung adenocarcinoma cell lines ([Barrettina et al., 2012](#)), which resulted in a similarly well-calibrated test statistic ([Figure S2E](#)).

Next, we tested for enrichment of loss-of-function (LOF) mutations by considering only truncating mutations as functional and all remaining mutation types as neutral. We applied this method to 2,266 genes with evidence of expression in lung adenocarcinoma cell lines and at least one truncating mutational event. Finally, we applied both PPH2 and LOF InVEx analyses to a focused set of Cancer Gene Census (CGC) genes expressed in lung adenocarcinoma and mutated or amplified in one or more tumor types.

Statistical Driver Analysis Yields Previously Reported and Novel Lung Adenocarcinoma Genes

The primary PPH2 InVEx analysis yielded 13 genes with statistical evidence of positive selection ($q < 0.25$) ([Table S3A](#)). These included lung adenocarcinoma genes with nonsynonymous mutation frequencies that were consistent with previous reports: *TP53* (50%), *KRAS* (27%), *EGFR* (17%), *STK11* (15%), *KEAP1* (12%), *NF1* (11%), *BRAF* (8%), and *SMAD4* (3%). This analysis also uncovered five novel candidates, including *CHEK2*, a gene driven by an apparent recurrent mapping artifact in three tumors and removed from all subsequent analyses (see [Extended Experimental Procedures](#)). The remaining candidates were mutated at frequencies lower than most previously reported genes, demonstrating the increased power of our large sample set. The LOF InVEx yielded six significantly mutated genes ($q < 0.25$), including *BRD3*, which is an additional gene not contained in the PPH2 analysis ([Table S3B](#)). The CGC-only PPH2 and LOF analyses yielded 15 and 10 genes, respectively, including *CTNNB1*, *FGFR3*, *ATM*, *CBL*, *PIK3CA*, *PTEN*, *FBXW7*, *ARID1A*, and *SETD2* ([Tables S3C](#) and [S3D](#)). In total, the union of these four analyses nominated 25 genes as significantly mutated in our cohort ([Figure 2A](#)). Somatic coding mutations in significantly mutated genes and known lung adenocarcinoma genes are provided in [Table S1](#). The entire list of somatic coding mutations for all covered genes is provided in [Table S4](#).

To compare our results with previous reports, we reviewed the CGC and lung adenocarcinoma literature to identify genes with previous evidence for functional somatic mutation in lung adenocarcinoma (see [Extended Experimental Procedures](#) for criteria and references). Of the 19 genes with reported functional mutations, 13 were significantly mutated genes nominated by our analysis (*KRAS*, *TP53*, *EGFR*, *STK11*, *SMARCA4*, *NF1*, *RB1*, *BRAF*, *KEAP1*, *SMAD4*, *CTNNB1*, *PIK3CA*, and *ATM*). The alter-

ations driving the statistical enrichment of these genes included previously reported and novel mutations ([Figures S3A–S3C](#)). The remaining six reported lung adenocarcinoma genes (*CDKN2A*, *ERBB2*, *AKT1*, *NRAS*, *HRAS*, and *APC*) were not significant in our mutation analysis ([Table S3E](#)), although we did identify canonical driver mutations in these genes (e.g., *AKT1* p.E17K, *NRAS* p.Q61L) and although *CDKN2A* is significantly deleted (see [Figure 2](#)) and rearranged (see below). This may reflect a power limitation of our cohort or analytic methods we applied, particularly when identifying infrequently mutated genes such as *AKT1*, *NRAS*, and *HRAS*. Also missing among our significantly mutated genes were 22 genes nominated by two previous large-scale targeted lung adenocarcinoma sequencing studies of similar or smaller size ([Ding et al., 2008](#); [Kan et al., 2010](#)) (see [Extended Experimental Procedures](#) for the complete list). Most of these genes (20 of 22) did not pass our gene expression filter and thus were not included in our global analysis. Targeted analysis of these genes identified four with nominal evidence for positive selection via PPH2 InVEx (*EPHA3*, *LPHN3*, *GRM1*, and *TLR4*), the most significant of these being *EPHA3* ($p = 0.0027$, PPH2 InVEx).

Correlations among Alterations in Significantly Mutated Genes and Clinicopathologic and Genomic Features

We correlated mutation status of the 25 significantly mutated genes with clinical features (smoking, age, and stage), genomic variables (mutation rate, mutation spectrum cluster, and imputed smoking status), and presence of driver alterations in 25 genes frequently or functionally altered in lung adenocarcinoma. These alterations included genes with reported high frequency of somatic mutation (e.g., *KRAS*) or focal amplification (e.g., *NKX2-1*) or deletion (e.g., *TP53*). High-frequency somatic copy number alterations used for this analysis were curated from published surveys of lung adenocarcinoma ([Tanaka et al., 2007](#); [Weir et al., 2007](#)). See [Hallmarks Analysis in the Experimental Procedures](#) for the strict definition of driver alterations. In our cohort, we observed gains of *TERT* (42% of cases, 15% focal), *MYC* (31%), *EGFR* (22%), and *NKX2-1* (18%, 10% focal). Frequent losses were seen in *TP53* (18%) and *CDKN2A* (24%, 10% homozygous), as well as in other significantly mutated genes, including *SMAD4*, *KEAP1*, and *SMARCA4*.

EGFR mutations were significantly anticorrelated with *KRAS* mutations ($p = 3.3 \times 10^{-4}$), and somatic mutation rate ($p = 5.9 \times 10^{-4}$) *EGFR* mutations significantly correlated with never-/light smoker status ($p = 2.0 \times 10^{-6}$), imputed never-/light smoker status (1.5×10^{-4}), and membership in spectrum cluster 1 ($p = 0.0015$). *KRAS*, *STK11*, *SMARCA4*, and *KEAP1* mutations were significantly anticorrelated with both spectrum cluster 1 and imputed never-/light smoking status ($p < 0.005$). These findings are consistent with reported associations ([Koivunen et al., 2008](#); [Pao et al., 2004, 2005](#); [Slebos et al., 1991](#)). In addition, *NF1* mutations were significantly depleted in spectrum cluster 1 ($p = 4 \times 10^{-3}$) and co-occurred with *U2AF1* mutations ($p = 0.0011$). *KRAS* driver alterations (including both mutations and copy number alterations) significantly associated with spectrum cluster 3 ($p = 0.00071$). *STK11* driver alterations were significantly enriched in spectrum cluster 2 ($p = 0.0026$). Correlation results are graphically summarized in [Figure S3D](#).

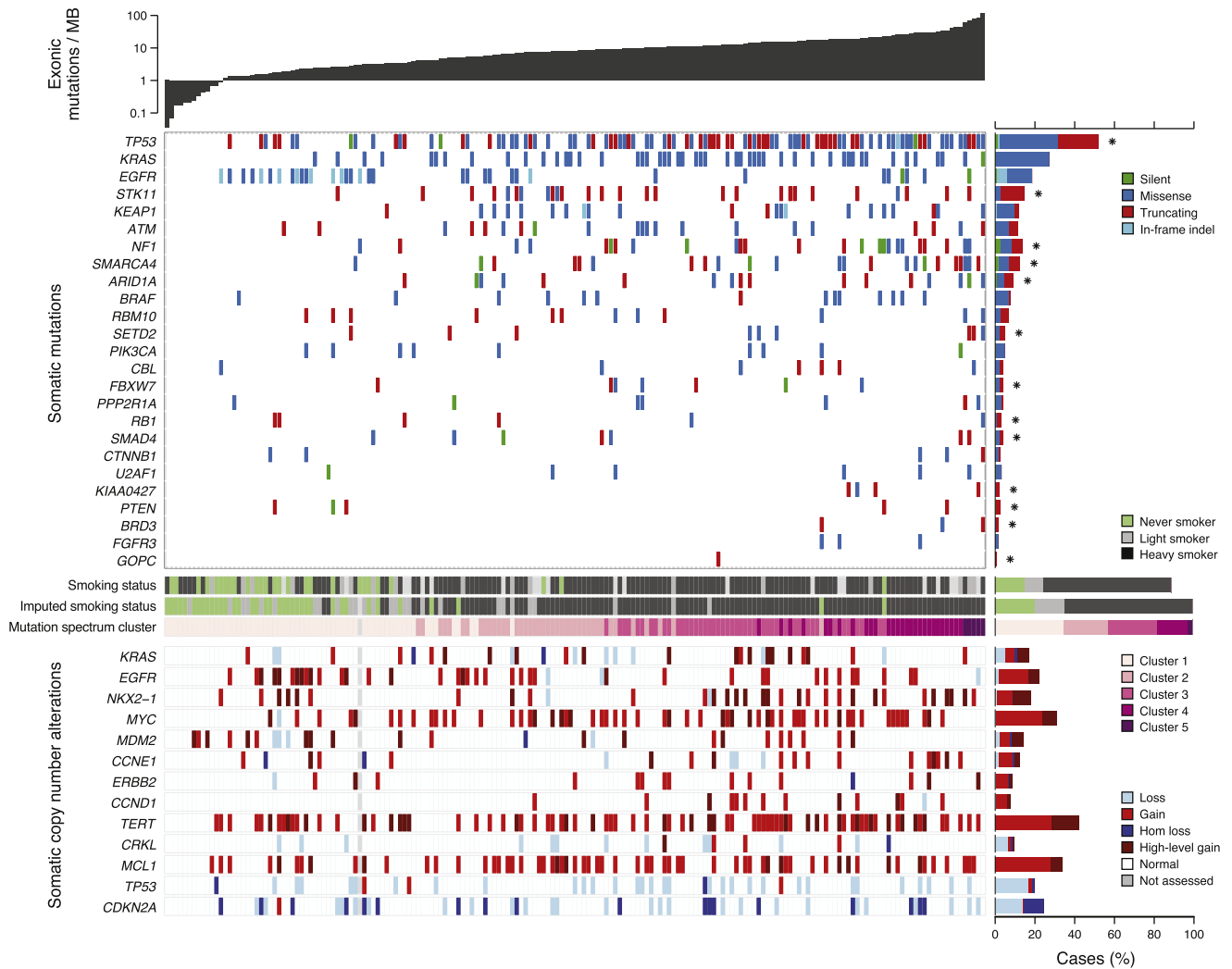


Figure 2. Somatic Mutations and Copy Number Changes in 183 Lung Adenocarcinomas

Top panel shows a summary of exonic somatic mutations in 25 significantly mutated genes (see text and Table S3 for details). Tumors are arranged from left to right by the number of nonsilent mutations per sample, shown in the top track. Significantly mutated genes are listed vertically in decreasing order of nonsilent mutation prevalence in the sequenced cohort. Colored rectangles indicate mutation category observed in a given gene and tumor. Bar chart (right) indicates prevalence of each mutation category in each gene. Asterisks indicate genes significantly enriched in truncating (nonsense, frameshift) mutations. Middle bars indicate smoking status and mutation spectrum cluster for each patient. White boxes indicate unknown status. Bottom panel shows a summary of somatic copy number alterations derived from SNP array data. Colored rectangles indicate the copy number change seen for a given gene and tumor. See also Figure S2.

Finally, we screened the 25 significantly mutated genes for association with progression-free survival (PFS) across 135 patients with PFS data. *U2AF1* ($p = 0.00011$, log rank test) and *TP53* mutations ($p = 0.0014$, log rank test) were associated with significantly reduced survival (Figure S3E). The latter finding was consistent with previous reports (Kosaka et al., 2009; Mitsudomi et al., 1993). No other significant associations with PFS were seen.

Nomination of Candidate Lung Adenocarcinoma Genes

One of the most significantly mutated genes in this lung adenocarcinoma cohort was *U2AF1* ($p = 2.0 \times 10^{-6}$, PPH2 lnVEx), which had nonsynonymous mutations in 3% of cases (Figure 3A). Identical c.101C > T, p.S34F mutations were seen in four of five *U2AF1* mutant cases (Figure 3A); this is the exact mutation re-

ported in myelodysplastic syndrome (MDS) (Graubert et al., 2012; Yoshida et al., 2011). To our knowledge, this study is the first report of *U2AF1* mutations in an epithelial tumor. One of four p.S34F mutations occurred with an activating event in *KRAS* (p.Q61H), suggesting that *U2AF1* mutations may confer tumorigenic capability independent of known proliferation-sustaining driver genes. As mentioned above, four patients with *U2AF1* mutations and survival data had significantly reduced PFS (Figure S3E). Nonsynonymous mutations in genes encoding other members of the spliceosome complex (including *SF3B1*, *U2AF2*, and *PRPF40B*) were found in 14 additional cases (Yoshida et al., 2011).

RBM10 was frequently mutated (12/183 cases; 7%) and subject to recurrent nonsense, frameshift, or splice-site mutations,

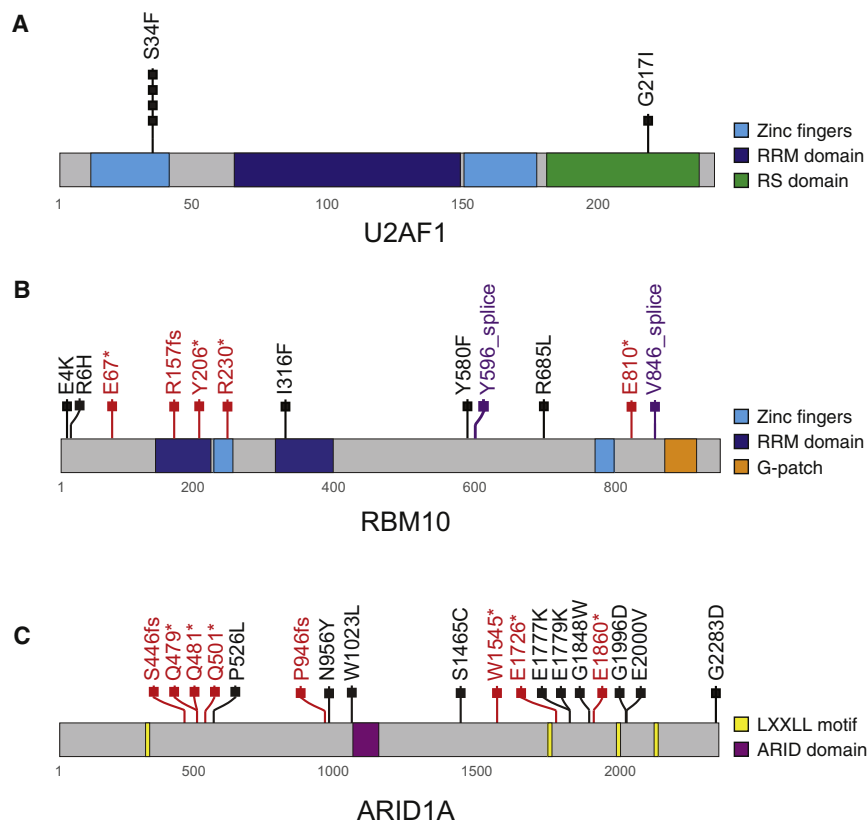


Figure 3. Somatic Mutations of Lung Adenocarcinoma Candidate Genes *U2AF1*, *RBM10*, and *ARID1A*

(A) Schematic representation of identified somatic mutations in *U2AF1* shown in the context of the known domain structure of the protein. Numbers refer to amino acid residues. Each rectangle corresponds to an independent, mutated tumor sample. Silent mutations are not shown. Missense mutations are shown in black.

(B) Schematic of somatic *RBM10* mutations. Splice-site mutations are shown in purple; truncating mutations are shown in red. Other notations as in (A).

(C) Schematic of somatic *ARID1A* mutations. Notations as in (A) and (B).

See also Figure S3.

which were present in 7 of 12 mutated cases (4% of overall cohort) (Figure 3B). This resulted in significant enrichment in the global PPH2 InVex analysis ($p = 0.00042$) (Table S3). Like *U2AF1*, *RBM10* is an RNA-binding protein that is highly expressed in lung adenocarcinoma cell lines (data not shown), and its mutations co-occurred with those in known lung adenocarcinoma oncogenes (*KRAS*, *EGFR*, and *PIK3CA*). *ARID1A*, encoding a key protein in the SWI/SNF chromatin-remodeling complex, was mutated in 8% of cases (Figure 3C) and showed significant accumulation of nonsense substitutions and frame-shift indels ($p = 0.027$, CGC LOF InVex).

Whole-Genome Rearrangement Analysis Reveals Novel and Recurrent Structural Variants

We used paired-end and split-read mapping of whole-genome data (Banerji et al., 2012; Bass et al., 2011; Medvedev et al., 2009) to detect and map the breakpoints of 2,349 somatic rearrangements across 24 WGS cases. The majority of these were intrachromosomal rearrangements (1,818 events) but included 531 interchromosomal events. Among these were 1,443 (61.4%) genic rearrangements (i.e., in which one breakpoint was contained within the promoter, UTR, intron, or exon of a gene) and 906 (38.6%) purely intergenic events. Lung adenocarcinomas harbored a wide range of total rearrangements (median: 98; range: 18–246), genic rearrangements (median: 50; range: 12–173) (Figure 4A), and overall genome complexity (Figure S4). The variability of rearrangement counts between cases did not

correlate with clinical variables (Figure S4 and Table S1) or mutation spectrum. Rearrangement coordinates and interpretations are provided as Table S5.

The reading frame of affected genes was preserved by 3% of detected rearrangements (71 of 2,349). These included 34 protein fusions, 13 duplications, and 24 deletions. We found 44 rearrangements that fused UTRs of two genes without affecting the protein-coding sequence of either gene. All 25 genic fusions we tested were confirmed by PCR and Illumina sequencing (see Extended Experimental Procedures) (Table S5).

The gene with the highest rate of rearrangements for its size was *CDKN2A* (4.3 rearrangements/sequenced Mb). Two cases had out-of-frame, antisense fusions (with *MTAP* and *C9orf53*), and a third harbored an in-frame deletion (Figure 4B). As shown in lung squamous cell carcinomas, rearrangements represent an additional mechanism of *CDKN2A* inactivation, in addition to reported mutation, homozygous deletion, and methylation (Cancer Genome Atlas Research Network, 2012). Additional lung adenocarcinoma tumor suppressors affected by predicted null or truncating rearrangements included *STK11* (2.5 kb deletion removing the translational start site) and *APC* (midexon rearrangement) (Figure 4B).

We next focused on potentially activating in-frame rearrangements of kinase genes. This analysis uncovered a two-exon deletion in *EGFR*, which was previously identified in glioblastoma multiforme but is novel in lung adenocarcinoma, ablating a portion of the C terminus of *EGFR* encoded by exons 25 and 26 (Figures 4B, 5A, and S5), including residues associated with interaction with *PIK3C2B* (Wheeler and Domin, 2001) and *CBL* (Grøvdal et al., 2004). Similar C-terminal deletion variants (*EGFR* vIvB) have been previously identified in glioblastoma (Ekstrand et al., 1992) and have been shown to be oncogenic in cellular and animal models (Cho et al., 2011; Pines et al., 2010). This tumor contained a second somatic alteration in *EGFR*, a p.G719S mutation, suggesting possible synergy of

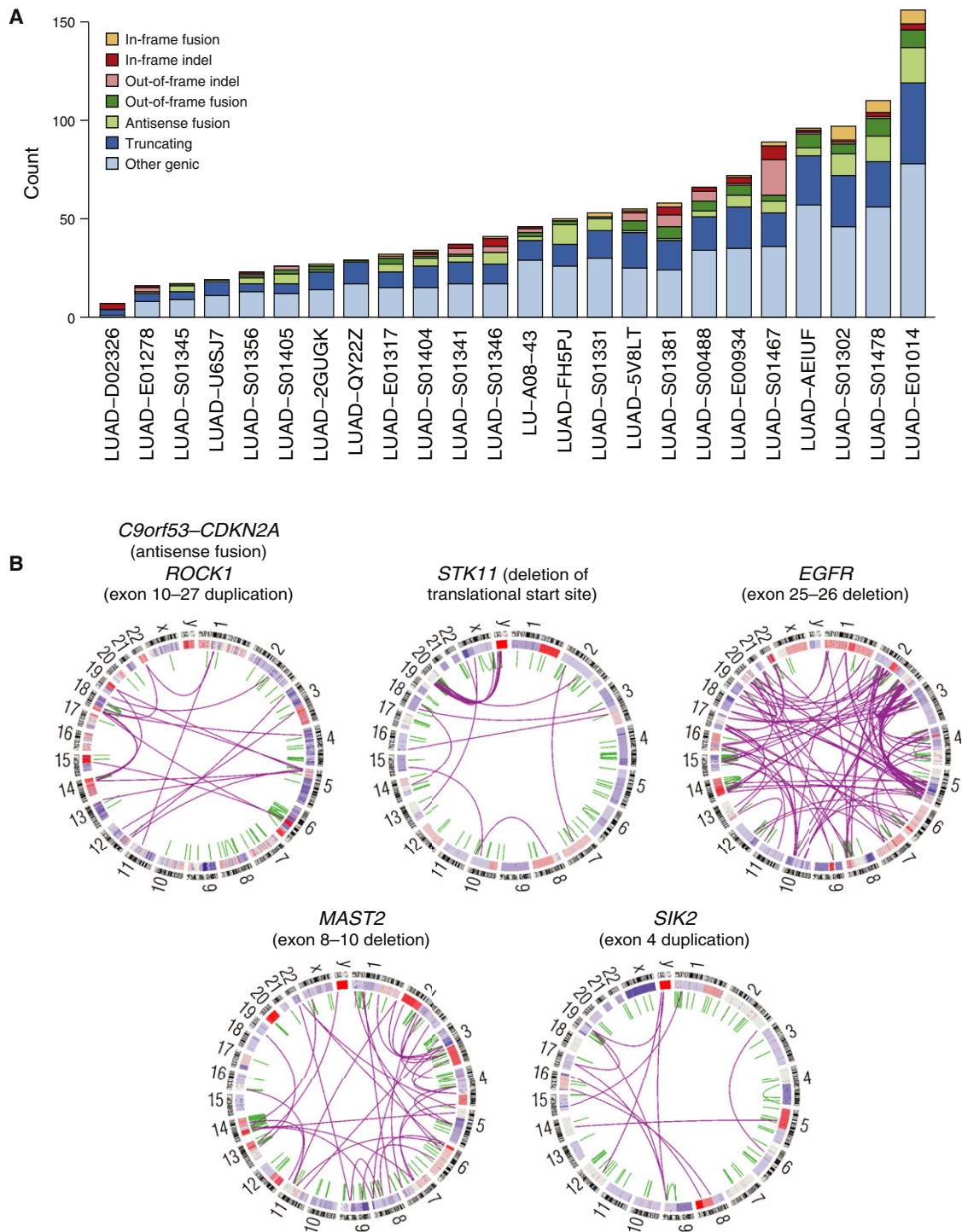


Figure 4. Whole-Genome Sequencing of Lung Adenocarcinoma

(A) Summary of genic rearrangement types across 25 lung adenocarcinoma whole genomes. Stacked-bar plot depicting the types of somatic rearrangement found in annotated genes by analysis of whole-genome sequence data from 25 tumor/normal pairs. The “Other Genic” category refers to rearrangements linking an intergenic region to the 3’ portion of a genic footprint.

(B) Representative Circos (Krzywinski et al., 2009) plots of whole-genome sequence data with rearrangements targeting known lung adenocarcinoma genes *CDKN2A*, *STK11*, and *EGFR* and novel genes *MAST2*, *SIK2*, and *ROCK1*. Chromosomes are arranged circularly end to end with each chromosome’s cytobands marked in the outer ring. The inner ring displays copy number data inferred from WGS with intrachromosomal events in green and interchromosomal translocations in purple.

See also Figure S4 and Table S5.

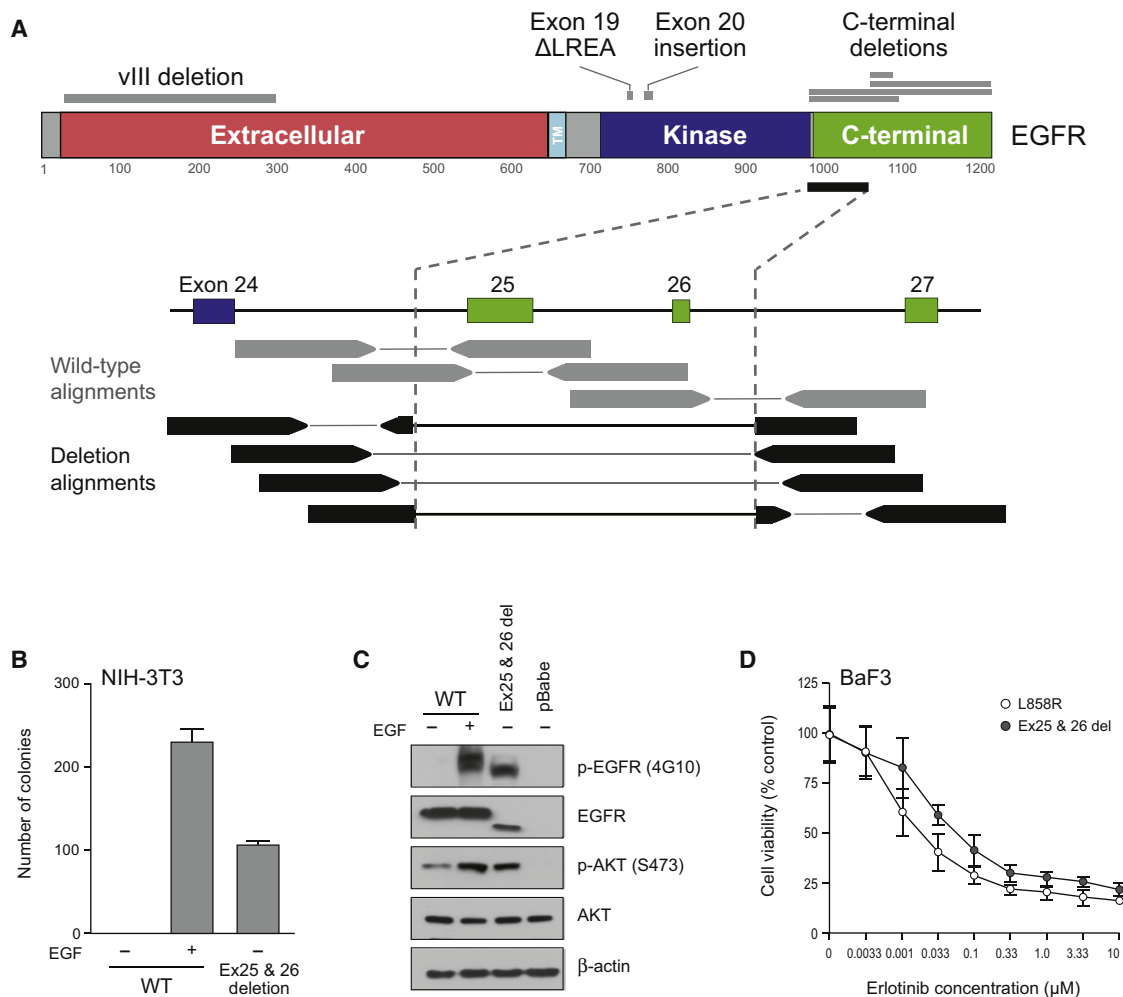


Figure 5. Identification of a Novel Lung Adenocarcinoma In-Frame Deletion in *EGFR*

(A) Schematic representation of reported *EGFR* alterations (above protein model) for comparison with a C-terminal deletion event found in this study by WGS (below protein model). A schematic depiction of sequencing data shows the expected wild-type reads (gray) in contrast with the observed reads (black) spanning or split by the deletion breakpoint. Supporting paired-end and split-read mapping data are shown in Figure S5.

(B) Soft agar colony forming assay of NIH 3T3 cells expressing exon 25- and 26-deleted *EGFR* (Ex25&26 del) or wild-type *EGFR* in the presence or absence of ligand stimulation. The bar graph shows the number of colonies formed by indicated cells with or without EGF in soft agar. Data shown are mean \pm SD of three replicates of a single experiment. The results are representative of three independent experiments.

(C) Ex25&26 del *EGFR* is constitutively active in the absence of EGF. The same NIH 3T3 cells used for the assay in (B) were subjected to immunoblotting with anti-phospho-tyrosine (4G10), anti-*EGFR*, and anti-phospho-Akt (S473) antibodies. Blots were probed with anti-Akt and anti- β -actin antibodies (loading control).

(D) Cell growth induced by the oncogenic *EGFR* deletion mutant is suppressed by erlotinib treatment. Ba/F3 cells transformed by either L858R or Ex25&26 del mutants were treated with increasing concentrations of erlotinib as indicated for 72 hr and were assayed for cell viability. Data shown are mean \pm SD of six replicates of a single experiment. The results are representative of three independent experiments.

See also Figure S5.

activating *EGFR* mutations or presence of independent, subclonal activating mutations.

To assess oncogenicity of this novel *EGFR* variant, we ectopically expressed an *EGFR* transgene lacking exons 25 and 26 in NIH 3T3 cells. As has been previously observed for oncogenic *EGFR* mutations, cells stably expressing this transgene demonstrated colony formation in soft agar (Figure 5B) and increased *EGFR* and AKT phosphorylation in the absence of EGF (Figure 5C). In contrast, cells expressing wild-type *EGFR* formed colonies only in the presence of EGF (Figure 5B). Overexpression

of the *EGFR* transgene in Ba/F3 cells led to interleukin-3 independent proliferation that was blocked by treatment with an *EGFR* tyrosine kinase inhibitor, erlotinib (Figure 5D), at concentrations previously shown to be sufficient for inhibition of activated variants of *EGFR* (Yuza et al., 2007).

Kinases with in-frame rearrangements in tumors without mutations in lung adenocarcinoma oncogenes included *SIK2* and *ROCK1* (Figure 4B). An in-frame kinase domain duplication in *SIK2* (salt-inducible kinase 2) was identified and validated by quantitative PCR (qPCR). The duplication occurred 15 amino

acids upstream of Thr-175, where a related kinase, SIK1, is activated by STK11 (Hashimoto et al., 2008). A 19 exon duplication was uncovered in *ROCK1*, which is a serine/threonine kinase that acts as an effector of Rho signaling (Pearce et al., 2010).

Notably, we did not identify any in-frame rearrangements involving kinase fusion targets in lung adenocarcinoma *ALK*, *RET1*, and *ROS1*. Given their reported 2%–7% frequency in lung adenocarcinoma (Bergethon et al., 2012; Takeuchi et al., 2012), our study of 24 tumor/normal pairs may not be large enough to detect these rearrangements. Interestingly, an out-of-frame *ROS1-CD74* translocation was identified in a single patient without evidence for the previously characterized reciprocal activating event. In-frame fusions and indels are annotated for each WGS case in Table S1.

DISCUSSION

Charting the Next-Generation Hallmarks of Lung Adenocarcinoma

The “hallmarks of cancer,” as defined by Hanahan and Weinberg (2000, 2011), comprise a set of cellular traits thought to be necessary for tumorigenesis. They also represent a powerful framework to evaluate our understanding of genetic alterations driving lung adenocarcinoma. With this aim, we mapped each of 25 experimentally validated lung adenocarcinoma genes to one or more cancer hallmarks from Hanahan and Weinberg (2000, 2011) (Table S6 and Experimental Procedures). These 25 genes include the 19 previously reported genes discussed above, in addition to six genes subject to frequent copy number alteration in lung adenocarcinoma (*NKX2-1*, *TERT*, *PTEN*, *MDM2*, *CCND1*, and *MYC*). Next, we integrated this gene hallmark mapping with our somatic mutation and copy number data to estimate the prevalence of cancer hallmark alterations in lung adenocarcinoma (Figure 6 and Table S1).

For many cases in our cohort, we could attribute only a minority of the ten cancer hallmarks to a distinct genetic lesion (Figure S6). Only 6% of tumors had alterations assigned to all six classic hallmarks, and none had alterations impacting all ten emerging and classic hallmarks. In contrast, 15% of our cohort did not have a single hallmark alteration, and 38% had three or fewer. This finding is likely explained in part by alteration of cancer genes by mechanisms not assayed in our study and also suggests that many lung adenocarcinoma genes have not been identified. This may be especially relevant for the hallmarks of avoiding immune destruction and tumor-promoting inflammation, to which none of the recurrently mutated genes identified in our study or previous studies could be linked. One of the most important and therapeutically targetable cancer hallmarks is sustaining proliferative signaling (Figures 6 and S6). Less than half (47%) of our cohort harbored a mutation in a known driver gene for this hallmark, and only slightly more (55%) did so when including high-level amplification in one or more proliferative signaling genes (e.g., *EGFR*, *ERBB2*, and *MYC*).

Our mapping of somatic alterations to cancer hallmarks illuminates specific gaps in the understanding of the somatic genetic underpinnings of lung adenocarcinoma. Around half of the sequenced cohort lacked a mutation supporting sustained proliferative signaling, and a majority lacked a genetic alteration

explaining the phenotypes of invasion and metastasis or angiogenesis. This phenotypic gap may be explained by novel capabilities not yet attributed to alterations in known lung adenocarcinoma genes or through novel alterations in genes previously unassociated with this disease that will emerge through additional unbiased analyses.

While annotating the 25 known lung adenocarcinoma genes, we noted that *SMARCA4*, an epigenetic regulator and tumor suppressor, could not be clearly mapped to any cancer hallmark. Given the frequent somatic mutations in epigenetic and splicing regulators found by recent cancer genome scans (Elsässer et al., 2011) and our study (*U2AF1*, *ARID1A*, *RBM10*, *SETD2*, and *BRD3*), we speculated that these alterations may represent a novel hallmark of epigenetic and RNA deregulation. Together, these genes implicate the proposed eleventh hallmark in a considerable proportion of cases (10% including only *SMARCA4*, 22% including nominated genes).

Efficiency and Power in Somatic Genetic Studies of Lung Adenocarcinoma

This study represents the largest sequencing analysis of lung adenocarcinoma to date. Our analysis reveals the genomic complexity of lung adenocarcinoma at the base-pair and structural levels, exceeding that observed in genome characterization studies of most other tumor types. We have applied a recently published statistical method (Hodis et al., 2012) for identifying somatically mutated genes displaying evidence of positive selection in cancer. This permutation approach exploits the abundant supply of intronic and flanking mutation events detected in both WES and WGS to adequately model the gene-specific variation in neutral mutation rates (Hodis et al., 2012). We believe that such a calibrated approach is required to identify signals of positive somatic selection in large unbiased cancer genome scans. This concern is particularly relevant to tumor types harboring high rates of somatic mutation, such as lung adenocarcinoma or melanoma.

This study has led to discovery of significant mutation of 25 genes in lung adenocarcinoma. Notably, our study did not identify a mutated oncogene in every tumor sample. Furthermore, we were unable to statistically nominate several important, but rarely mutated, lung adenocarcinoma genes (*AKT1*, *ERBB2*, *NRAS*, and *HRAS*, each with ≤ 3 events in our cohort). Therefore, future studies of larger cohorts by The Cancer Genome Atlas and other consortia that combine analysis of data from RNA sequencing (RNA-seq), methylation profiling, and other omic platforms will likely yield an even more complete annotation of genes significant to lung adenocarcinoma.

Conclusion

This study represents a significant advance toward complete characterization of the genomic alterations of lung adenocarcinoma. These results are a testament to the power of unbiased, large-scale next-generation sequencing technology to expand our understanding of tumor biology. The novel mutated genes identified in this study warrant further investigation to determine their biologic, prognostic, and/or therapeutic significance in lung adenocarcinoma, potentially leading to clinical translation and improved outcomes for patients with this deadly disease.

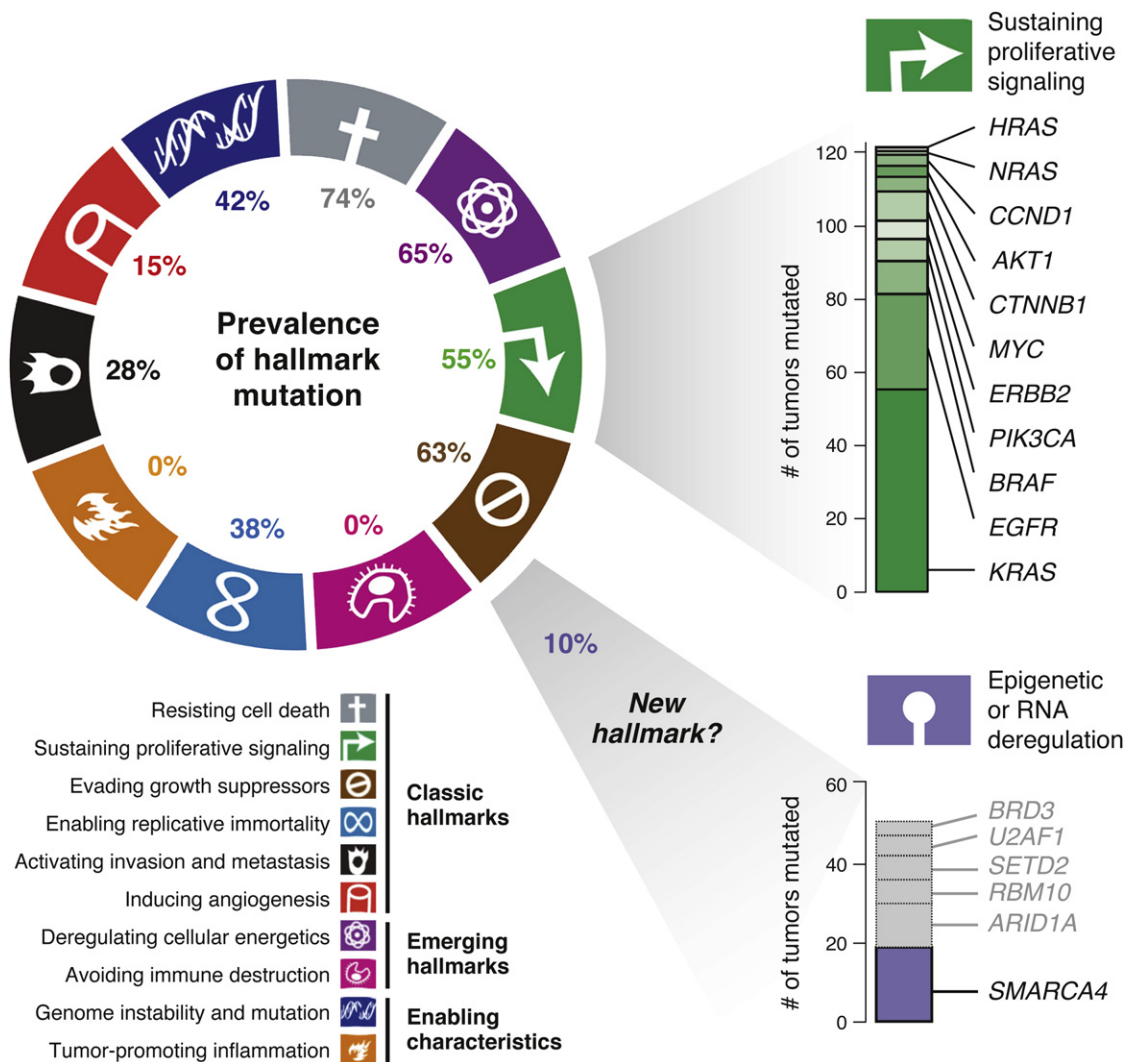


Figure 6. Next-Generation Hallmarks of Lung Adenocarcinoma

Left, the prevalence of mutation or SCNA of Sanger Cancer Gene Census (Futreal et al., 2004) genes mapping to cancer hallmarks defined by Hanahan and Weinberg (2011). Suspected passenger mutations were filtered out of the analysis, as described in Experimental Procedures. Top right, genes comprising the mutated genes in the hallmark of sustaining proliferative signaling are shown, depicted as above. Genes shown in gray are candidate lung adenocarcinoma genes identified in this study that may additionally contribute to the hallmark.

See also Figure S6 and Table S6.

EXPERIMENTAL PROCEDURES

Details of sample preparation and analysis are described in the Extended Experimental Procedures.

Patient and Sample Characteristics

We obtained DNA from tumor and matched normal adjacent tissue from six source sites. DNA was obtained from frozen tissue primary lung cancer resection specimens for all samples, with the exception of one patient (LU-A08-14), for whom a liver metastasis was obtained at autopsy. The 183 lung adenocarcinoma diagnoses were either certified by a clinical surgical pathology report provided by the external tissue bank or collaborator or was verified through in-house review by an anatomical pathologist at the Broad Institute of MIT and Harvard. A second round of pathology review was conducted by an expert committee led by W.D.T. Informed consent (Institutional Review Board) was

obtained for each sample by using protocols approved by the Broad Institute of Harvard and MIT and each originating tissue source site.

Massively Parallel Sequencing

Exome capture was performed by using Agilent SureSelect Human All Exon 50 Mb according to the manufacturer’s instructions. All WES and WGS was performed on the Illumina HiSeq platform. Basic alignment and sequence quality control were done by using the Picard and Firehose pipelines at the Broad Institute. Mapped genomes were processed by the Broad Firehose pipeline to perform additional quality control, variant calling, and mutational significance analysis.

External Data

Gene expression data for 40 lung adenocarcinoma cell lines were obtained from the Cancer Cell Line Encyclopedia (CCLE) (<http://www.broadinstitute>).

org/ccl/home) as RMA normalized tab-delimited text data (Barretina et al., 2012).

Statistical Analyses

We evaluated statistical evidence for somatic selection within the longest transcript of each gene by using InVEx (Hodis et al., 2012) with PolyPhen-2-based (Adzhubei et al., 2010) and LOF-based scoring schemes. The method was implemented in Python (<http://www.python.org>) and is available for download (<http://www.broadinstitute.org/software/invex/>). Gene ranking according to a stratified binomial model was performed by using the MutSig method from Berger et al. (2011) and was implemented in MATLAB. Correlations between genotype status, mutation/rearrangement spectrum data, and clinical variables were performed by Fisher's exact test for dichotomous variables and by Wilcoxon rank sum test for dichotomous variables versus numeric data (e.g., mutation status versus total mutation rate). All remaining statistical computing, including cluster analysis and visualization, was performed by using standard packages in R (<http://www.r-project.org>).

Hallmarks Analysis

We manually assigned 25 genes—implicated by previous studies to be frequently or functionally altered in lung adenocarcinoma—to one or more cancer hallmarks as defined by Hanahan and Weinberg (2000, 2011) (see Extended Experimental Procedures). We determined whether alterations in gene *i* could be implicated as a “driver” of one or more cancer hallmarks in case *j* by applying the following criteria: we inferred the activation status of genes annotated by the Sanger Gene Census as “dominant” cancer genes (e.g., *KRAS*) in each patient by evaluating every nonsynonymous variant in the gene for its presence within a COSMIC hot spot (Forbes et al., 2011). Mutations that were present in the COSMIC database (<http://www.sanger.ac.uk/genetics/CGP/cosmic/>) at least ten times were considered oncogenic mutations. We considered a dominant gene activated if it harbored such a variant or a high-level, focal amplification. We considered recessive cancer genes (e.g., *TP53*) to be inactivated if the gene had (1) a truncating mutation, (2) compound missense mutations, (3) a hemizygous missense mutation, or (4) homozygous copy number loss. We mapped each patient *j* to hallmark *k* if the sample contained at least one activating or inactivating event in a dominant or recessive cancer gene, respectively, that mapped to hallmark *k*.

ACCESSION NUMBERS

The dbGAP accession number for the data reported in this paper is phs000488.v1.p1.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Extended Experimental Procedures, six figures, and six tables and can be found with this article online at <http://dx.doi.org/10.1016/j.cell.2012.08.029>.

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