Non-Small Cell Lung Cancer Radiogenomics Map Identifies Relationships between Molecular and Imaging Phenotypes with Prognostic Implications¹

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Purpose:

To create a radiogenomic map linking computed tomographic (CT) image features and gene expression profiles generated by RNA sequencing for patients with non-small cell lung cancer (NSCLC).

Materials and Methods:

A cohort of 113 patients with NSCLC diagnosed between April 2008 and September 2014 who had preoperative CT data and tumor tissue available was studied. For each tumor, a thoracic radiologist recorded 87 semantic image features, selected to reflect radiologic characteristics of nodule shape, margin, texture, tumor environment, and overall lung characteristics. Next, total RNA was extracted from the tissue and analyzed with RNA sequencing technology. Ten highly coexpressed gene clusters, termed metagenes, were identified, validated in publicly available gene-expression cohorts, and correlated with prognosis. Next, a radiogenomics map was built that linked semantic image features to metagenes by using the t statistic and the Spearman correlation metric with multiple testing correction.

Results:

RNA sequencing analysis resulted in 10 metagenes that capture a variety of molecular pathways, including the epidermal growth factor (EGF) pathway. A radiogenomic map was created with 32 statistically significant correlations between semantic image features and metagenes. For example, nodule attenuation and margins are associated with the late cell-cycle genes, and a metagene that represents the EGF pathway was significantly correlated with the presence of ground-glass opacity and irregular nodules or nodules with poorly defined margins.

Conclusion:

Radiogenomic analysis of NSCLC showed multiple associations between semantic image features and metagenes that represented canonical molecular pathways, and it can result in noninvasive identification of molecular properties of NSCLC.

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Online supplemental material is available for this article.

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Study supported by National Cancer Institute (R01CA160251) and National Institute of Biomedical Imaging and Bioengineering (R01EB020527).

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ORIGINAL RESEARCH - THORACIC IMAGING

on-small cell lung cancer (NSCLC) is the most common type of lung cancer and is composed of tumors with significant molecular heterogeneity resulting from differences in intrinsic oncogenic signaling pathways (1). Molecular characteristics of NSCLC formed the basis of clinical lesion diagnosis and therapeutic treatment (2,3). For example, the activation of the epidermal growth factor (EGF) receptor pathway determines treatment with anti-EGF receptor therapy by using tyrosine kinase inhibitors (4). Such molecular properties of NSCLC were recently characterized (5) by quantitative imaging signatures. These findings and similar results in other cancers (6-8) confirm the potential synergy of integrating imaging and genomic data (9), yielding insights into understanding lesion-specific features and transcriptional regulators in patients with NSCLC.

Recent development of RNA sequencing techniques presents new opportunities for characterization of molecular pathways in NSCLC (10,11). Accordingly, the emergence of radiogenomics (7,12,13) allows for identification of noninvasive biomarkers, which reflects cellular and molecular properties of NSCLC. These noninvasive

Advances in Knowledge

- Ten molecularly defined metagenes had 32 significant associations with CT image features in patients with non-small cell lung cancer (NSCLC) (false discovery rate, <0.01).
- Radiologist-observed CT characteristics that captured nodule attenuation and nodule margins were associated with the late cell-cycle genes (false discovery rate, <0.01).
- Radiologist-observed CT characteristics of the degree of ground-glass opacity and poorly defined and irregular margins of lung nodules were associated with the epidermal growth factor molecular pathway (false discovery rate, <0.01).

imaging biomarkers act as surrogates for molecularly defined features and may enable noninvasive precision medicine. Our radiogenomics analysis has several advantages over previous studies (Fig 1) (14,15). First, for the study of each patient, we collected an extensive collection of semantic features that consisted of 87 features defined by using a controlled vocabulary and that reflected radiologic characteristics of lung nodules (eg, nodule location, shape and texture of the tumor, and features derived from the lesion macroenvironment such as presence and patterns of emphysema and fibrosis). Second, we used RNA sequencing to characterize the transcriptomic profile of each tumor. We summarized these data by defining metagenes as clusters of coexpressed genes and used gene-enrichment analysis to annotate these metagenes with distinct molecular pathways. Additionally, we studied molecular prognostic significance by measuring overall survival predictors from public cohorts, which showed the prognostic associations of each metagene in two important NSCLC histologic structures: adenocarcinoma and squamous cell carcinoma. The purpose of our study was to create a radiogenomic map that linked features from computed tomographic (CT) images and gene expression profiles generated by RNA sequencing for patients with NSCLC.

Materials and Methods

Image Data Collection and Annotation

With institutional review board approval, we studied 113 patients who underwent surgery for NSCLC between April 2008 and September 2014 at two

Implication for Patient Care

 Multiple characteristics of lung nodules observed at CT by radiologists were associated with distinct molecular pathways in NSCLC, which may help guide clinicians in noninvasive treatment planning. medical centers and who underwent pretreatment chest CT examination and had tissue available for RNA sequencing (Table 1). Image data were obtained from GE medical systems (Waukesha, Wis) and Siemens (Erlangen, Germany) scanners. CT section thickness was as follows: less than 1 mm (22 patients, 19.5%), 1 mm to less than 2 mm (88 patients, 77.9%), and 2-3 mm (three patients, 2.6%). We developed a lung annotation template to facilitate selection of up to 87 semantic image features (Table E1 [online]) by using the open-source ePAD platform (https:// epad.stanford.edu) that enables quantitative imaging annotations (16). These features reflected radiologic observations that included nodule shape, margin, texture, and location, and overall lung characteristics. A thoracic radiologist (A.L., with 20 years of chest oncologic imaging experience) annotated the CT image of each tumor by using ePAD while blinded to all clinical and molecular information. The semantic image features have binary values that reflect presence (or absence) of the radiologic features except for several variables that are ordinal in nature. These include nodule features that describe nodule margin (ie, smooth, irregular, lobulated, spiculated, and poorly defined), nodule shape (four classes from

https://doi.org/10.1148/radiol.2017161845

Content codes: CT CH

Radiology 2018; 286:307-315

Abbreviations:

EGF = epidermal growth factor NSCLC = non-small cell lung cancer

Author contributions:

Guarantors of integrity of entire study, M.Z., 0.G.; study concepts/study design or data acquisition or data analysis/interpretation, all authors; manuscript drafting or manuscript revision for important intellectual content, all authors; approval of final version of submitted manuscript, all authors; agrees to ensure any questions related to the work are appropriately resolved, all authors; literature research, M.Z., A.L., S.N., 0.G.; clinical studies, M.Z., A.L., J.B.S., G.J.B.; experimental studies, M.Z., S.E., S.N., 0.G.; statistical analysis, M.Z., A.G., 0.G.; and manuscript editing, all authors

Conflicts of interest are listed at the end of this article.

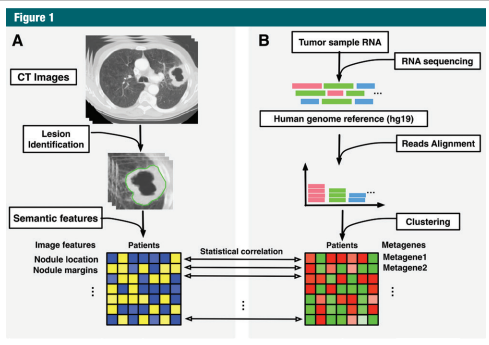


Figure 1: Overview of radiogenomic analysis to identify associations between, *A*, semantic features at CT and, *B*, RNA sequencing data.

round to polygonal), nodule attenuation (four classes from pure solid to ground glass), and nodule ground-glass composition (6° from 0%–100%). We also evaluated for the presence, type, location, and distribution of emphysema and fibrosis. All features are reported in Table E1 (online). To remove features with less variant frequencies, we chose to study semantic features with occurrence rate of greater than 10% in the study cohort, which resulted in the removal of 52 features.

Next-generation RNA Sequencing Protocol

The tumor samples were frozen after surgery at -80°C until use. They were obtained intraoperatively as the specimen was removed from the patient. A slice of tumor cut from the center of the longest diameter was harvested (depending on the overall dimensions of the tumor, this slice thickness varied from several millimeters to nearly 1 cm). This process was completed within 30 minutes or less to minimize time before samples were frozen. For larger tumors we avoided any areas of obvious central necrosis. Tumor purity was verified for approximately one

of every 10 lung tumors to be at least 50% surface area of a selected sample. Total RNA was extracted from nodule tissue samples and converted into a standard library of TruSeq Illumina kit for paired-end sequencing (Centrilion Biosciences, Palo Alto, Calif). The protocol included Ribo-Zero ribosomal RNA removal and ribosomal RNA depletion step, followed by fragmentation and complementary DNA with DNA synthesis by using SuperScript II (Life Technologies, Carlsbad, Calif). Quality was confirmed by using the BioAnalyzer (Agilent Technologies, Santa Clara, Calif), and the concentration was then evaluated by Kapa qPCR (Kapa Biosystems, Wilmington, Mass). RNA sequencing interpretations were mapped to the Human Genome version 19 (http://genome.ucsc.edu) by using the alignment algorithm (Star version https://github.com/alexdobin/ STAR). Next we used software (Cufflinks version 2.0.2; http://cole-trapnelllab.github.io/cufflinks/) to determine the expression calls in each sample by using the number of fragments per kilobase of transcript per million mapped interpretations. Values of fragments per kilobase of transcript per million mapped interpretations were subsequently log transformed, and missing values were estimated by using a 15-nearest-neighbors algorithm. We only included transcripts with at least five interpretations mapped to their location in at least 70% of the samples.

Statistical Analysis

By focusing on highly expressed gene expression data, we created metagenes with coherent gene expression as previously defined in Gevaert et al (15). We chose to select the top 10 metagenes with the highest cluster homogeneities in external gene expression data sets. We calculated the homogeneity score of each metagene by averaging all pairwise Pearson correlation coefficients of genes within the metagene and within each external gene expression data set. We used five public lung cancer geneexpression cohorts: the combined lung adenocarcinoma and squamous cell carcinoma from the Cancer Genome Atlas (17), the cohort from Lee et al (18), the cohort from Bild et al (19), the cohort from Shedden et al (20), and the cohort from Roepman et al

Characteristic	Result
Characteristic	nesuit
Average age (y)	69 (46–85)
Sex	
Male	86 (76.1)
Female	27 (23.9)
Histologic structure	
Adenocarcinoma	84 (74.3)
Squamous cell carcinoma	26 (23.0)
Other NSCLC	3 (2.7)
Smoking status	
Nonsmoker	16 (14.1)
Former smoker	74 (65.5)
Current smoker	23 (20.4)
EGFR	
Positive	17 (15.0)
Negative	73 (64.6)
Missing	23 (20.4)
KRAS	
Positive	21 (18.6)
Negative	69 (61.1)
Missing	23 (20.3)
No. of tumors according to stage $$	
Stage 1	
Adenocarcinoma	41
Squamous cell carcinoma	13
Other	1
Stage 2	
Adenocarcinoma	33
Squamous cell carcinoma	9
Other NSCLC	0
Stage 3	
Adenocarcinoma	7
Squamous cell carcinoma	3
Other NSCLC	1
Stage 4	
Adenocarcinoma	3
Squamous cell carcinoma	1
Other NSCLC	1

Note.—Unless otnerwise indicated, data are number or patients and data in parentheses are percentages. EGFR epidermal growth factor receptor, KRAS = V-Ki-ras2 Kirsten rat sarcoma viral oncogene homolog, Adeno = adenocarcinoma, SCC = squamous cell carcinoma. *Data in parentheses are range.

(21), for a total of 1227 patients with NSCLC. Next, we annotated the top 10 metagenes (Table 2) by using functional enrichment analysis of the metagenes (22). Briefly, we use the *P* value generated by a hypergeometric test that assessed whether the overlap between genes in a functional category and genes

Metagene Cha			•	-					
A: Metagene Char	acteristics Inc	luding N	umber of Gene	s, Functional A	nnotatio	ns, and k	Key Genes		
Top 10 Metagene	No. of Genes	Key Pa	athway Genes			Function	nal Annotation		
Metagene 19	24	LRIG1	, HPGD, GDF1:	5		EGF sur	vival pathway		
Metagene 10	45	CDH2	CDH2, POSTN, VCAN, PDGFRA			Extracellular matrix			
Metagene 9	38	VCAM	1, CD44, CD48	3		Immune response			
Metagene 4	133	CD4, LYL1, SPI1, CD37			T-cell activation				
Metagene 3	103	VIM, LM02, EGR2			Hypoxia and inflammation				
Metagene 21	38	BGN, COL4A1, COL5A1, COL5A2 Mesenchymal stem co			hymal stem cells	;			
Metagene 65	20	MUC1, TMEM101, TMEM125, TMEM41B Epithelial mesenchymal t				transitio			
Metagene 56	26	JUN, TXNIP, ZNF91, KLF2 Cellular			Cellular	r metabolism			
Metagene 60	25	EGR1,	EGR1, CD55, CXCL2			Mesothelioma survival signature			
Metagene 38	57	MCM ²	MCM4, MCM5, AURKA, AURKB, E2F2			Late cell division cycle			
B: The Homogene	ous Scores of	the Curr	ent Cohort and	Five Public Da	atasets				
	Current	TCGA	Lee et al	Bild et al	Shedd	en et al	Roepman et al		
Top 10 Metagene		TCGA Cohort	Lee et al Cohort (18)	Bild et al Cohort (19)	Shedd Cohor		Roepman et al Cohort (21)	Averag	
Top 10 Metagene Metagene 19	Cohort							Averag	
	Cohort 0.60	Cohort	Cohort (18)	Cohort (19)	Cohor		Cohort (21)		
Metagene 19	Cohort 0.60 0.58	Cohort 0.61	Cohort (18) 0.64	Cohort (19) 0.61	Cohori		Cohort (21) 0.53	0.57	
Metagene 19 Metagene 10	Cohort 0.60 0.58 0.56	Cohort 0.61 0.69	Cohort (18) 0.64 0.66	Cohort (19) 0.61 0.61	0.45 0.64		Cohort (21) 0.53 0.53	0.63	
Metagene 19 Metagene 10 Metagene 9	Cohort 0.60 0.58 0.56 0.56	Cohort 0.61 0.69 0.56	Cohort (18) 0.64 0.66 0.61	Cohort (19) 0.61 0.61 0.60	0.45 0.64 0.58		Cohort (21) 0.53 0.53 0.44	0.57 0.63 0.56	

Note.—In part A, the homogeneous scores of the cohort and five public datasets were also recorded; metagenes were ranked in a decreasing order in terms of their homogeneity scores. In part B, metagenes were ranked in a decreasing order in terms of their homogeneity scores. The last column in part B shows the average records of the five validation cohorts. TCGA = the Cancer Genome Atlas.

0.51

0.57

0.56

0.64

0.46

0.38

0.34

0.61

0.47

0.31

0.45

0.41

0.53

0.48

0.50

0.60

in a metagene was larger than expected by chance. We used the following databases that define functional categories for genes: MSigDB version 3 (23), GeneSetDB version 4 (24), Chea for Chip-X gene sets version 2 (25), and manually curated gene sets related to stem cells and immune gene sets. Gene set enrichment P values were corrected for multiple testing by using the false discovery rate (26), and we used a P value threshold less than .001 and Q-value threshold smaller than 0.05 to call a gene functional category significant.

0.49

0.46

0.46

0.45

0.57

0.54

0.54

0.65

0.66

0.59

0.58

0.66

Metagene 65

Metagene 56

Metagene 60

Metagene 38

We used PRECOG (https://pre-cog.stanford.edu/) (27), a tool that links genes with prognosis by using the largest collection of gene expression data to date, to assess the survival relationship of the top 10 metagenes in

adenocarcinoma and squamous cell carcinoma separately. PRECOG contains independent, publicly available geneexpression cohorts with survival data for lung adenocarcinoma (n = 17) and lung squamous cell carcinoma (n = 15). Within each metagene, the included genes were standardized to have no mean and unit variance in each cohort separately. For each dataset, association with overall survival was assessed by using univariate Cox proportional hazards regression (survival package version 2.38 in R version 3.1.2; R Foundation, Vienna, Austria). We reported the z score of each cohort to represent the direction and strength of the correlation of each metagene with overall survival (negative values indicated good prognosis and positive values indicated poor prognosis). Next, we computed a global meta-z score for each metagene by combining the z scores obtained from each public cohort by using the Stouffer method (27). A z score larger than 2 or smaller than -2 was considered to be statistically significant.

We built a radiogenomics map by associating metagenes with semantic image features. We used the *t* statistic and Spearman correlation metric to assess significant associations and used the false discovery rate (26) to correct for multiple testing. A *P* value less than .05 and a false discovery rate value less than 0.01 were used to determine statistically significant associations between metagenes and image features. We also reported image feature-to-metagene correlation coefficients to indicate directional relationship of metagenes and semantic features at CT imaging.

Results

Radiogenomics NSCLC Cohort

Table 1 shows the clinical characteristics of our cohort of 113 patients with NSCLC. For each tumor, we used the previously described template (28) to annotate 87 semantic image features that represented the radiographic phenotype of each tumor (Table E1 [onlinel). Removal of low-variance features resulted in 35 semantic image features that captured nodule location, nodule margins, nodule attenuation, nodule ground-glass composition, and the presence of emphysema for subsequent statistical analysis. Next, we used highthroughput RNA sequencing to capture the transcriptome information of these cases. This RNA sequencing process resulted in quantification of 60498 genes per sample represented by the ensemble identifiers, where transcripts that had at least five interpretations in 70% of the samples were included (see Materials and Methods section).

Identification of Coexpressed Metagenes

We created the previously defined 56 metagenes in NSCLC by using the RNA sequencing data collected here (9). Next, we validated the metagene

Table 3

Prognostic Performance of Metagenes Assessed By Meta Z Scores Summarizing the Individual Z Scores from PRECOG Analysis and Corresponding Meta P Value

	Adenoc	arcinoma	Squamous Cell Carcinoma			
Top 10 Metagene	Meta Z Score	FDR-corrected Meta <i>P</i> Value	Meta Z Score	FDR-corrected Meta <i>P</i> Value		
Metagene 19	-6.43	.006	-1.51	.786		
Metagene 10	0.29	.912	-0.45	.968		
Metagene 9	-3.95	.006	-3.08	.020		
Metagene 4	-0.72	.912	-2.28	.184		
Metagene 3	0.11	.912	-2.26	.184		
Metagene 21	-0.5	.912	-0.82	.968		
Metagene 65	-6.54	.006	0.28	.968		
Metagene 56	-5.23	.006	0.88	.968		
Metagene 60	-2.7	.035	0.04	.968		
Metagene 38	6.96	.006	1.83	.469		

Note.—Negative z scores indicate good prognosis and positive z scores indicate poor prognosis. See Table E2 (online) for detailed z scores for each individual adeno and squamous cell carcinoma data set. FDR = false discovery rate.

homogeneity in five external public cohorts for validation (see Materials and Methods section). Table 2b reports the metagene homogeneity in our cohort and the validation cohorts (metagenes were ranked in a decreasing order in terms of their homogeneity scores). A high value of homogeneity score indicates a high level of gene similarities within each metagene. We selected the top 10 metagenes with high average homogeneity (homogeneity score, ≥ 0.45) on the basis of their average score in validation cohorts for further analysis (Table 2b).

Metagene Functional Enrichment and Prognostic Assessment

We annotated molecular functions of metagenes by using gene set enrichment analysis (23), which allows identification of shared common biologic pathways from a public molecular signature database. We demonstrated that the top 10 metagenes captured a variety of known molecular classes including EGF pathway (29), genes related to the extracellular matrix (30), immune response (31), and T-cell activation (32) (Table 2a). Next, we used survival meta-analysis (27) to assess the survival relationship of metagenes and public genomic cohorts with survival outcomes (Tables 3, E2 [online]).

We showed that prognostic performance varied in groups of adenocarcinoma and squamous cell carcinoma. As shown in Table 3, five metagenes are significantly correlated with overall survival in adenocarcinoma data sets. For example, metagene 38 enriched by late cell division cycle genes is strongly correlated with poor prognosis, and four metagenes (metagenes 9, 19, 56, and 65) are correlated with positive survival. In squamous cell carcinomas, survival tended to be more heterogeneous with only three metagenes (metagenes 3, 4, and 9) and weakly correlated with good survival (Table 3).

Radiogenomics Map of NSCLC Revealed Associations

We built a radiogenomics map by correlating each image feature to the identified metagenes (Fig 2). We found 32 statistically significant correlations between semantic features and metagenes (Table 4; false discovery rate, <0.01). For example, we found that the metagene capturing the late cell cycle was associated with nodule attenuation and nodule margins. When this metagene was active, the lesion tended to be solid (metagene 38; false discovery rate, 0.01), whereas when this metagene was inactive, the lesion tended to have poorly defined margins

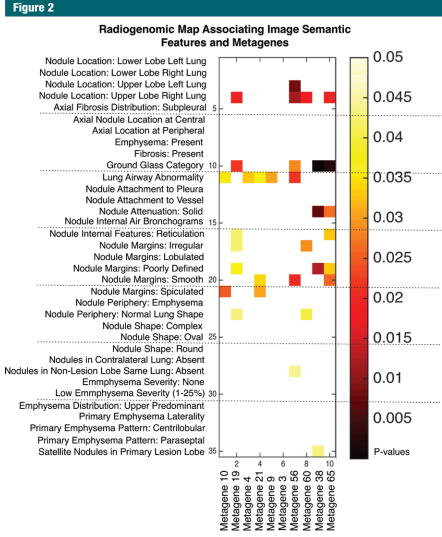


Figure 2: Radiogenomic map revealing 32 statistically significant associations between 35 CT semantic features and top 10 metagenes in NSCLC. Only image features above the variance cutoff of 10% are shown. The order of image features and metagenes is determined by hierarchical clustering of the *P* values.

(metagene 38; false discovery rate, 0.008). We also found that a normal lung background was positively correlated to a mesothelioma survival signature (metagene 60: false discovery rate, 0.008), which indicated that genes active in this pathway were associated with underlying lung parenchymal abnormalities such as emphysema when underexpressed and normal lung parenchymal morphologic structure when overexpressed. Next, metagene 65 was enriched with genes associated the epithelial-to-mesenchymal transition and had several significant

associations with image features (Table 4). For example, overexpression of metagene 65 was associated with poorly defined margins, and underexpression of metagene 65 was associated with smooth margins. Overall, our radiogenomic analysis captured multiple associations between semantic image features at CT and molecular pathway activity in NSCLC (Fig 2, Table 4).

EGF Pathway Characterizes Specific Images Phenotype of NSCLC

Next, we focused more deeply on the associations of metagene 19 with semantic features. Metagene 19 reflects the activity of the EGF pathway, which captures the relationship between the gene LRIG1 as a negative regulator of EGF. When metagene 19 is active, LRIG1 results in inhibition of cancer cell growth (33-35). Our radiogenomic map (Table 4) showed that metagene 19 was significantly associated with six semantic features at CT. Metagene 19 was negatively correlated with solid lesions and smooth margins (Fig 3a), which showed that low activity of metagene 19 was associated with solid lesions with smooth, lobulated, or spiculated margins. Conversely, high expression of metagene 19 was correlated with a high proportion of ground-glass opacity, irregular or poorly defined nodule margins, and reticulation (Fig 3b). More specifically, rising expression of metagene 19 was strongly correlated with nodule margin shape with increasing irregularity (Fig 4). Interestingly, this group of NSCLC was enriched in cases with EGF receptor mutations (P < .0001, Wilcoxon rank-sum test).

Discussion

In this study, we integrated semantic image features at CT with next-generation RNA sequencing data to identify radiogenomic biomarkers of NSCLC. RNA sequencing analysis revealed 10 metagenes that resulted in 32 significant pair-wise associations between quantitative image features and metagenes annotated by functional gene enrichment analysis. These associations show the feasibility of noninvasive molecular characterization of NSCLC by using radiogenomics mapping. We thoroughly validated the 10 metagenes in publicly available data sets by using their homogeneity, which reflected that coexpression is not only present in the cohort presented here, but also in five additional cohorts that represent 1227 patients from other institutes (Table 2). Moreover, correlation of the metagenes with clinical outcome in publicly available cohorts from patients from PRECOG (27) provided strong evidence of their prognostic significance

					Spearma
					Correlation
Parameter	Image Feature	<i>P</i> Value	FDR	t Statistic	Coefficie
Two-sample <i>t</i> test					
Metagene 10	Nodule margin: spiculated	.034	0.008	-2.148	
Metagene 10	Lung airway abnormality		0.009	2.336	
Metagene 19	Anatomic location: upper lobe of right lung	.015	0.009	-2.469	
Metagene 19	Nodule internal features: reticulation	.049	0.008	1.983	
Metagene 19	Nodule margin: irregular	.049	0.008	1.983	
Metagene 19	Nodule margin: poorly defined	.035	0.007	2.139	
Metagene 19	Nodule periphery: normal lung shape	.046	0.009	2.013	
Metagene 4	Lung airway abnormality	.031	0.008	-2.185	
Metagene 21	Lung airway abnormality	.034	0.007	-2.138	
Metagene 21	Nodule margin: smooth	.032	0.007	-2.168	
Metagene 21	Nodule margin: spiculated	.027	0.008	2.229	
Metagene 9	Lung airway abnormality	.028	0.008	-2.228	
Metagene 56	Anatomic location: upper lobe of left lung	.006	0.009	2.774	
Metagene 56	Anatomic location: upper lobe of right lung	.008	0.01	-2.665	
Metagene 56	Lung airway abnormality	.017	0.008	-2.420	
Metagene 56	Nodule margin: smooth	.015	0.008	-2.464	
Metagene 56	Nodules in nonlesion lobe in same lung: absent	.048	0.009	1.998	
Metagene 60	Anatomic location: upper lobe of right lung	.013	0.011	-2.506	
Metagene 60	Nodule margin: irregular	.026	0.008	2.257	
Metagene 60	Nodule periphery: normal lung shape	.042	0.008	2.059	
Metagene 38	Nodule attenuation: solid	.005	0.01	2.849	
Metagene 38	Nodule margin: poorly defined	.009	0.008	-2.645	
Metagene 38	Satellite nodules in primary lesion lobe	.046	0.009	-2.014	
Metagene 65	Anatomic location: upper lobe of right lung	.015	0.01	-2.469	
Metagene 65	Nodule attenuation: solid	.022	0.008	-2.310	
Metagene 65	Nodule internal features: reticulation	.031	0.008	2.188	
Metagene 65	Nodule margin: poorly defined	.031	0.008	2.187	
Metagene 65	Nodule margin: smooth	.021	0.008	-2.331	
Spearman correlation coefficient					
Metagene 19	Ground-glass category	.017	0.008		0.223
Metagene 56	Ground-glass category	.025	0.008		0.211
Metagene 38	Ground-glass category	<.001	0.03		-0.322
Metagene 65	Ground-glass category	.003	0.007		0.279

in independently collected gene-expression cohorts (Table 3).

Linking imaging characteristics with molecular signatures is a growing field of research that provides additional value to clinical imaging with relevant molecular biology information (Table 4). For example, metagene 19 and its image-feature associations are a prototypical example of the possibilities of radiogenomics mapping (Fig 3).

Low activity of this metagene reflects NSCLC lesions that trigger the EGF receptor pathway, which consequently activates KRAS and PIK3CA genes and results in cell proliferation. We showed that these tumors are characterized by smooth margins and appear to be solid as defined by high attenuation at CT. However, high activity of metagene 19 corresponds to activation of LRIG1, a negative inhibitor of EGF that results

in turning off the EGF receptor pathway. When these tumors are enriched in mutations in EGF receptor, thereby severing the inhibitory link with LRIG1, downstream activation of KRAS and PIK3CA pathways again occur but manifest with a different phenotype at CT as cancers with markedly irregular or poorly defined margins, most likely caused by the presence of ground-glass opacity. Overall, this example highlights that image phenotypes reflect different activities of molecular pathways and allow for noninvasive assessment of the molecular activity of NSCLC lesions with potential implications for treatment. Moreover, we can speculate that the radiogenomic map can be extended to capture therapy response of existing or novel agents through the use of gene signatures predicting the response of treatment. These signatures can be mapped to image features by using radiogenomics mapping, as discussed here through metagenes, and subsequently allow for noninvasive assessment of treatment management.

Our study has the following limitations. Because the collected cohort of patients with NSCLC included various section thicknesses and other acquisition parameters, future studies should determine the effects of scanner heterogeneity on the semantic annotations of radiologists. In addition, one thoracic radiologist annotated all semantic features for the study cohort. Future work that incorporates annotations by multiple radiologists is needed to study any potential variability in semantic feature annotation. Also, because this was a prospective cohort, direct survival analysis of the patients was not included because of lack of sufficient follow-up data of patients. To counter this, we introduced public datasets to associate metagenes with prognosis. We also opted to build a radiogenomics map in the largest possible cohort of NSCLC. Therefore, we focused on all NSCLCs, including adenocarcinoma and squamous cell carcinoma. Although the histopathologic classification is readily distinguishable in tissue samples, it is not always apparent from the imaging phenotype.

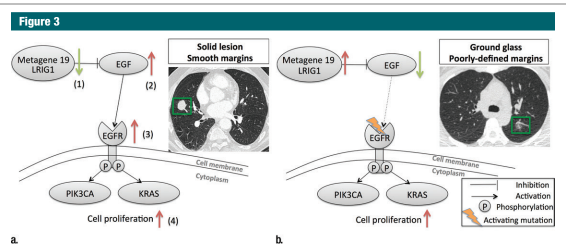


Figure 3: Association of the activity of metagene 19 with two distinct image phenotypes. (a) Low activity of metagene 19 is associated with low activity of LRIG1 (green arrow), and high activity of the EGF–EGF receptor (*EGFR*; red arrow) pathway results in cell proliferation through activating KRAS and PIK3CA. (b) High activity of metagene 19 results in high activity of LRIG1 (red arrow) with inhibition of the EGF–EGF receptor pathway (green arrow), but results in higher occurrence of EGF receptor mutations, which severs the link between LRIG1 and EGF receptor.

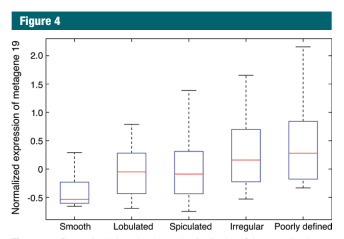


Figure 4: Box-and-whisker plot shows the distribution of the normalized expression of metagene 19 regarding different types of nodule margins at CT. The normalized expression of metagene 19 reflects the activity of the genes in this metagene.

In summary, in this study we presented a radiogenomics map of NSCLC that linked image phenotypes with RNA signatures captured by metagenes and how they are associated with molecular pathways. This extensive radiogenomics map allowed for a better understanding of the pathophysiologic structure of lung cancer and how molecular processes manifest in a macromolecular way as captured by semantic image features. The presented image-to-molecular

feature associations open possibilities for assessing therapeutic options on the basis of biologic pathway activity by using surrogate image features. Moreover, adding other molecular measurements such as DNA methylation or DNA copy number can deepen the radiogenomics associations and increase the potential of building radiogenomics maps even further, and enable a noninvasive in-depth understanding of lung cancer biology by using CT images.

Acknowledgment: We are grateful to Dr Denise Aberle, professor of Radiology the University of California Los Angeles for her invaluable input to develop the semantic template.

Disclosures of Conflicts of Interest: M.Z. disclosed no relevant relationships. A.L. disclosed no relevant relationships. S.E. disclosed no relevant relationships. A.G. disclosed no relevant relationships. J.B.S. Activities related to the present article: disclosed no relevant relationships. Activities not related to the present article: author reports grants and personal fees from Varian, personal fees from Carefusion (Benton-Dickinson), and personal fees from Maquet, outside the submitted work. Other relationships: disclosed no relevant relationships. K.C.J. disclosed no relevant relationships. G.J.B. disclosed no relevant relationships. S.K.P. Activities related to the present article: disclosed no relevant relationships. Activities not related to the present article: author disclosed a pending patent for systems, methods, and devices for analyzing quantitative information obtained from radiologic images. Other relationships: disclosed no relevant relationships. D.L.R. disclosed no relevant relationships. S.N. Activities related to the present article: disclosed no relevant relationships. Activities not related to the present article: author disclosed board membership for Fovia and consultancy for Carestream. Other relationships: disclosed no relevant relationships. O.G. disclosed no relevant relationships.

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