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Success and failure rates of tumor genotyping techniques in routine pathological samples with non-small-cell lung cancer

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Abstract

Introduction—Identification of some somatic molecular alterations in non-small-cell lung cancer (NSCLC) has become evidence-based practice. The success and failure rate of using commercially-available tumor genotyping techniques in routine day-to-day NSCLC pathology samples is not well described. We sought to evaluate the success and failure rate of *EGFR* mutation, *KRAS* mutation, and *ALK* FISH in a cohort of lung cancers subjected to routine clinical tumor genotype.

Methods—Clinicopathologic data, tumor genotype success and failure rates were retrospectively compiled and analyzed from 381 patient-tumor samples.

Results—From these 381 patients with lung cancer, the mean age was 65 years, 61.2% were women, 75.9% were white, 27.8% were never smokers, 73.8% had advanced NSCLC and 86.1% had adenocarcinoma histology. The tumor tissue was obtained from surgical specimens in 48.8%, core needle biopsies in 17.9%, and as cell blocks from aspirates or fluid in 33.3% of cases. Anatomic sites for tissue collection included lung (49.3%), lymph nodes (22.3%), pleura (11.8%), bone (6.0%), brain (6.0%), among others. The overall success rate for *EGFR* mutation analysis was 94.2%, for *KRAS* mutation 91.6% and for *ALK* FISH 91.6%. The highest failure rates were observed when the tissue was obtained from image-guided percutaneous transthoracic core-needle biopsies (31.8%, 27.3%, and 35.3% for *EGFR*, *KRAS*, and *ALK* tests, respectively) and bone specimens (23.1%, 15.4%, and 23.1%, respectively). In specimens obtained from bone, the failure

CONFLICT OF INTEREST STATEMENT

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rates were significantly higher for biopsies than resection specimens (40% vs 0%, p=0.024 for EGFR) and for decalcified compared to non-decalcified samples (60% vs 5.5%, p=0.021 for EGFR).

Conclusions—Tumor genotype techniques are feasible in most samples, outside small imageguided percutaneous transthoracic core-needle biopsies and bone samples from core biopsies with decalcification, and therefore expansion of routine tumor genotype into the care of patients with NSCLC may not require special tissue acquisition or manipulation.

Keywords

lung cancer; non-small-cell lung cancer; never smokers; epidermal growth factor receptor; EGFR; anaplastic lymphoma kinase; ALK; KRAS; tumor genotype; failure; bone specimen; core biopsy; molecular testing

INTRODUCTION

In the United States in 2013, the expected number of new cases of and deaths from lung cancer will exceed 220,000 and 159,000, respectively (1). The overall, of all stages combined, five-year survival for the most prevalent form of lung cancer - non-small-cell lung cancer (NSCLC) - does not exceed 15% despite use of surgical resection, radiotherapy and systemic chemotherapy (1). The last decade of research in lung cancer has yielded important advances in the development of targeted therapies that target driver oncogenes (2). The most prevalent mutated or rearranged oncogenes identified in non-small cell lung cancers (NSCLCs) are v-ki-ras2 Kirsten rat sarcoma viral oncogene homolog (KRAS), epidermal growth factor receptor (EGFR), anaplastic lymphoma kinase (ALK), ROS1, BRAF, ERBB2 and RET (3). Specifically, mutations in EGFR and rearrangements (either inversions or translocations) involving ALK are part of the pathogenesis of some lung adenocarcinomas, predominantly in never and/or light smokers, and predict for improved outcomes with tyrosine kinase inhibitors (TKIs), such as erlotinib and crizotinib, respectively, that target these aberrant kinases (4–6). As such, molecular testing of lung cancer specimens has become part of routine clinical practice in the care for patients with advanced NSCLC.

Currently, the proposed testing guidelines from the College of American Pathologists, the International Association for the Study of Lung Cancer, the Association for Molecular Pathology, and the National Comprehensive Cancer Network recommend testing all advanced NSCLCs with an adenocarcinoma component at the time of diagnosis for *EGFR* mutation and for *ALK* fluorescence in situ hybridization (FISH) analysis (7; 8). These specimens are more frequently encountered as small biopsy or cytology specimens derived from either the primary tumor or from lymph node or distant metastatic sites, and processed as formalin-fixed, paraffin embedded tissue samples.

Limited tumor cellularity in small biopsy or cytology specimens can lead to molecular testing failure. Additionally, it is recognized that different tissue processing techniques, including acid decalcification or heavy metal fixatives, can cause DNA degradation and impede molecular analysis (9). These issues can influence the selection and success rates of specimens submitted for mutational analysis when considering sampling of bony lesions. Practically, this can pose a dilemma for the clinician and pathologist alike, in determining whether an initial biopsy used to establish the diagnosis of NSCLC is sufficient for additional molecular testing, or whether another specimen must be procured for such a purpose.

As molecular testing of NSCLCs has become a standard of clinical practice, various institutions have reported their testing protocol and experience (10–13). From these studies, one can occasionally glean the failure rates of their specimens, though to our knowledge a systematic study of genetic testing failure rates for NSCLC using commercially-available testing in a typical clinical practice setting has not been published to date. Here, we present the molecular testing efficiency for *EGFR*, *KRAS*, and *ALK* FISH analysis of clinical specimens from NSCLC patients from our institution over a five year time period, reflecting real-life clinical practice experience.

MATERIALS AND METHODS

Patient selection

Patients with a diagnosis of lung cancer who were seen by our providers and whose tumors were genotyped for at least *EGFR* mutations were identified through an ongoing Institutional Review Board (IRB) approved protocol at Beth Israel Deaconess Medical Center (BIDMC2009-P-000182). Patients and tumor pairs were excluded if genotyping was not performed. There were 381 patient-tumor specimens that were submitted to a commercial vendor for tumor genotype techniques between 2007 and 2012. The data cut off for analyses was December 19th, 2012. Study data were collected and managed using REDCap electronic data capture tools hosted at BIDMC.

Tumor processing and genotype

Surgical (i.e., either incisional or excisional biopsies that required a surgical procedure) and core needle biopsies were processed using standard techniques: 10% neutral buffered formalin fixation and paraffin-embedding. For any bone specimens that could not be cut with a scalpel at the grossing bench,, an acid decalcification was performed using RDO rapid decalcifier solution (Apex Engineering Products Corporation, Aurora, Illinois) following formalin fixation. Core needle and small biopsy specimens typically underwent a brief 15-30 minute decalcification, whereas larger surgical resection specimens were decalcified for 2-6 hours depending on the amount of calcified bone in the chosen sections. Cell aspirates or cell concentrates from fluid samples were collected into a methanol-water fixative (CytoLyt, Hologic Corp., Marlborough, MA) and a single ThinPrep slide prepared, with residual material used to create a cell block using a plasma-thrombin method prior to formalin-fixation and paraffin embedding. Once a diagnosis was established on histologic and/or immunohistiologic staining profiles as per evidence-based recommendations (8), the residual material in the formalin-fixed paraffin-embedded (FFPE) tissue blocks were submitted for molecular analysis. When multiple tissue blocks were available, the one with the highest tumor cellularity was chosen, without additional tumor microdissection or enrichment. Molecular analysis of tumor specimens was performed by a commercial vendor, Integrated Oncology (LabCorp, Esoterix Genetic Laboratories, LLC). EGFR mutation analysis was performed using standard DNA sequencing techniques with exons 18 to 21 sequenced (14; 15). For KRAS mutation analysis, DNA from exon 2 was amplified and subjected to single nucleotide primer extension to detect mutations at codons 12 and 13. ALK translocation status was analyzed using the Vysis ALK Break-Apart fluorescence in situ hybridization (FISH) probe (Abbott Molecular, Inc., Des Plaines, IL), as previously described (16). Failure of the assays was defined as insufficient/unusable material to isolate DNA or inability to perform/complete sequencing for EGFR and KRAS mutations, and lack of hybridization signals after two attempts for ALK FISH.

Data collection

Clinical, pathologic, radiographic and tumor genotyping information was collected from chart extraction. The site of biopsy (lung, lymph node, pleura, bone, brain, liver,

pericardium, or adrenal) and the type of biopsy (surgical specimen [both excisional and incisional], core needle biopsy, or cell block from aspirate/fluid) were extracted from the medical record. Slides from all specimens that failed molecular testing, as well as a subset of the successfully genotyped cases were re-reviewed by a pathologist (PAV), with data compiled on tumor cellularity, use of ancillary studies, histopathologic features, and specimen processing including decalcification.

Statistical methods

Logistic regression was used for the univariable and multivariable analyses to capture the influence of each clinical predictive factor on the failure rate of tumor genotyping. The clinically relevant predictive factors were included into the multivariable analysis regardless of their statistical significance in the univariable analysis. All categorical variables were dichotomized in the regression analyses. We report odds ratio (OR) and 95% confidence interval (95%CI) to each predictive factor. Fisher's exact test was performed to compare categorical variables. *p*-value < 0.05 was considered as statistically significant. All *p*-values we reported were two-sided. We performed our statistical analyses with STATA version 12 (STATA Corp, College Station, TX).

RESULTS

Patient and tumor characteristics

Table 1 summarizes the clinical and pathological characteristics of the 381 patient-tumor pairs that were included in our cohort.

Success and failure rates of tumor genotype techniques

Out of the 381 samples, all were sent for *EGFR* mutation analysis, 226 for *KRAS* mutation analysis, and 275 for *ALK* FISH (Table 1). In 207 tumors (54.3%), all three tests were ordered simultaneously. Table 2 details the success and failure rates for *EGFR*, *KRAS*, and *ALK* analyses techniques. The success rate for *EGFR* mutation analysis was 94.2% and the failure rate 5.8%; for *KRAS* mutation 91.6% and 8.36%, respectively; and for *ALK* FISH 91.6% and 8.41%, respectively. In the 207 tumors in which the three tests were ordered, the success rate for *EGFR* was 92.3%, for *KRAS* 91.8% and for *ALK* FISH 89.9% (Table 2). Out of the 25 failure cases of the latter cohort, 11 failed *EGFR*, *KRAS* and *ALK* concurrently, 4 failed *ALK* FISH alone, 3 each failed *EGFR/ALK* or *KRAS/ALK* pairs, 2 failed *KRAS* alone, and 1 *EGFR/KRAS* or *EGFR* mutation analysis alone

Tumor characteristics that correlate with increased failure rate

To determine the possible clinical or tumor characteristics that were associated with an increased incidence of failure of tumor genotyping techniques, we performed both univariable and multivariable logistic regressions in the 381 samples and focused on *EGFR* mutation analysis (due to its higher number of patient-tumor samples). Table 3 indicates the results. The patient's sex, ethnicity, smoking history, tumor stage, and histologic subtype were not associated with an increase in failure of *EGFR* mutation analysis. However, nonsurgical forms of tumor acquisition (core needle biopsies or cytologic specimens with cell blocks), with an odds ratio (OR) of 28.45 (p=0.002), and bone as the site of tumor acquisition, with an OR of 6.16 (p=0.010), were significantly associated with increase failure rates for *EGFR* mutational analysis (Table 3). Similarly, non-surgical forms of tumor acquisition, with an OR of 4.32 (p=0.006), and bone as the site of tumor acquisition, with an OR of 3.58, p=0.03, were significantly associated with increase failure rates for *ALK* FISH analysis (data not shown).

To better define these tumor characteristics associated with genotype failure, we analyzed the cohort for differences in success rate by site and type of tissue acquisition (Tables 4 and 5).

Type of tissue acquisition and failure rate of tumor genotype

Out of the 207 patients with concurrent testing, the highest failure rates were observed when the tissue was obtained from core needle biopsies (30.8%, 20.5%, and 30.8% for *EGFR*, *KRAS*, and *ALK* tests, respectively) when compared to larger surgical resection specimens (0%, 4.21%, and 3.16% for *EGFR*, *KRAS* and *ALK* tests, respectively) or cell block preparations (5.47%, 8.21% and 6.84% for *EGFR*, *KRAS* and *ALK* tests, respectively. In the latter group, cell block preparations from aspirates or fluid had similar failure rates [data not shown]).

We further on determined that out of the core needle biopsies, the failure rates were higher when the tissue of origin was either bone or lung when compared to all other sites (Table 5).

In particular, the failure rate of image-guided percutaneous transthoracic core-needle biopsies was high at 31.8% for *EGFR* mutations while the failure rate for bronchoscopy-guided lung transbronchial biopsies was much lower at 11.1% (Table 5). We compared the two aforementioned types of lung biopsies in relation to estimated tumor cellularity, tumor area, use of touch preparation for rapid on-site cytologic evaluation, presence of extensive desmoplastic stromal response, and number of slides cut from the paraffin block used for diagnostic immunohistochemical and ancillary studies. The most significant differences observed in the retrievable specimens were in the use of a touch preparation (image-guided percutaneous transthoracic core-needle biopsies 17/19 cases versus 0/15 cases for bronchoscopy-guided lung transbronchial biopsies, p<0.0001) and the number of slides obtained (image-guided percutaneous transthoracic core-needle biopsies had 9/14 cases with 10 slides versus 2/12 cases for bronchoscopy-guided lung transbronchial biopsies, p=0.0214). These findings may infer increased tumor tissue use or loss in image-guided

Bone specimens and failure of tumor genotype

Table 5 shows the success and failure rate by site of tumor acquisition. The highest failure rates for tumors with more than ten specimens sent for molecular analysis were observed in samples derived from bone (Tables 4 and 5). The failure rate for *EGFR* mutation analysis of tumors obtained from bone was significantly higher than in all other sites combined (4/23 vs. 18/340, p=0.0357). Out of the 207 patients with concurrent testing, the failure rate for bone-derived specimens was 23.1%, 15.4%, and 23.1% for *EGFR*, *KRAS*, and *ALK* tests, respectively (data not shown).

percutaneous transthoracic core-needle biopsies prior to tumor genotyping.

We further went on to examine factors that might explain the higher failure rate in specimens originating from bone. The failure rate was significantly different based on the type of tissue acquisition, with a higher failure rate from non-surgical biopsy or cytologic cell block specimens from bone (Table 4). The failure rate was significantly higher in non-surgical biopsies versus surgical resections for *EGFR* mutations (40% [4/10] vs. 0% [0/13], respectively, p=0.024), *ALK* FISH (44.4% [4/9] vs. 0% [0/9], p=0.082) and *KRAS* mutations (2/9 [22.2%] vs. 0% [0/5], p=0.505).

In addition, decalcification of the tumor tissue prior to preparation of the paraffin sample correlated with a higher rate of genotype failure. For *EGFR* mutation analysis, the failure rate for decalcified versus non-decalcified samples was significantly higher (60% [3/5] vs. 5.5% [1/18], respectively, p=0.021). Representative examples of the histologic features of

bone-derived specimens that either were successful for or failed subsequent genotyping are provided in Figure 1.

DISCUSSION

We have shown that our institutional success rate for EGFR, KRAS, and ALK testing of clinical lung cancer specimens exceeds 90% using routine tissue samples and a commercial vendor. As molecular testing of non-squamous, non-small cell lung carcinomas has become the standard of care for patients with advanced stage lung cancer, numerous studies by highvolume academic centers have been published outlining mutation frequency in different patient populations (17). Less has been published regarding the actual testing success or failure of tumor specimens in clinical practice. Overall tumor genotyping success rates extracted from various studies demonstrate a range from 81-94% following initial pathologist screen for sufficient tumor cell in the specimen (10; 13; 18; 19). Comparable results have also been described more specifically for small biopsy and cytology-derived specimens, with success rates ranging from 83-100% in published studies (12; 20; 21). A large proportion of these studies perform molecular/FISH testing in-house, whereas here we describe a large cohort of patient samples with molecular/FISH testing performed by an independent commercial lab. It is encouraging to find that the overall molecular testing success rate of 90-94%, as well as the success rate for small biopsy or cytology derived specimens of 85–88%, demonstrated in this study are right in line with the reported testing success rates in the literature. Thus, comparable success rates can be achieved in a send-out manner for institutions or medical centers that either do not have in-house molecular testing capabilities, or otherwise choose to outsource such tests.

After identifying the overall tumor genotyping success rates; we performed a detailed analysis using multivariate statistics to identify those factors that might predict testing failure, and found that while clinical characteristics (patient sex, ethnicity, smoking status, tumor stage, histologic subtype) had no impact on testing failure, the method of tumor tissue acquisition and the anatomic site from which that tumor came did correlate with tumor testing success rates. The finding that smaller biopsy or cytology-derived specimens had a higher chance of testing failure than surgical specimens is clearly linked to the amount of tumor cells obtained for analysis: larger specimens generally have ample tissue for diagnostic workup and testing, whereas the tumor tissue present in small biopsy/cytology specimens may be largely used for the diagnostic work-up limiting the material available for subsequent testing. Along these lines, recommendations for limited immunohistochemical work-up and tissue conservation strategies have been put forth to both maximize the diagnostic accuracy as well as preserve as much tissue as possible for molecular testing in these small lung cancer specimens (22).

The majority of tumor genotyping failures from lung specimens were from image-guided percutaneous transthoracic core-needle biopsies. These biopsies had a high frequency of rapid on-site cytologic evaluation in the form of touch-imprint cytologic slides prepared from the core biopsies. This method of cytologic on-site assessment transfers tumor cells from the tissue core to the cytologic slide, in doing so reduces the number of tumor cells that remain in the core that is used for diagnosis and molecular testing. Additionally, it was found that more slides on average were cut from the tissue blocks of image-guided percutaneous transthoracic core-needle biopsies than other forms of lung biopsies (such as transbronchial biopsies). More slides cut from the paraffin block mean less material remaining in the block for subsequent testing. When considering other institution's experience, a recent large-scale study for the feasibility of image-guided percutaneous transthoracic core-needle biopsies for lung tumor genotyping reports slightly lower testing

failure rates (approximately 20%), though it appears as if cytologic on-site evaluation was not utilized (20).

Tumor cells obtained from bony metastasis is the other specimen characteristic that was associated with a higher rate of molecular testing failure. The routine processing of these specimens incorporating an acid decalcification step so that histologic sections can be cut poses problems for downstream molecular testing. Acid decalcification of bone tissue extensively fragments DNA and also interferes with subsequent FISH analysis (9), and should be avoided if possible for metastatic lung cancer specimens. Even when ample tumor cellularity is present, acid decalcification often causes subsequent molecular testing failure (Figure 1A). If the amount of calcified bone tissue is small and the specimen can be cut without a decalcification step, one greatly increases the chances of successful molecular testing (Figure 1B). Along these lines, a clinically or radiologically characterized "bone lesion" can in reality have little or no bone tissue in the biopsy specimen and obviates the need for tissue decalcification (Figure 1C). Therefore, it is important for pathologists and clinicians alike to recognize that just because a specimen is labeled as a bone biopsy or a bony lesion, this should not preclude molecular testing.

From the clinical standpoint, there are many factors that go into deciding what putative tumor site to biopsy, and how to obtain tissue for diagnosis and ancillary testing. For patients that initially present with clinically or radiologically apparent stage IV disease, the principal goal in obtaining a tissue diagnosis should be to both diagnosis and pathologically stage the patient's lung cancer simultaneously, using the safest, least invasive, and least costly test available (23). Although open surgical biopsies are the gold standard for obtaining ample diagnostic tissue, the associated morbidity/mortality and cost that can be associated with this modality has motivated the shift towards less invasive small biopsy or fine needle aspiration of lesions in establishing a pathologic diagnosis of cancer. As has been shown in this report as well as others, the use of these minimally invasive small biopsy or cytologic-derived specimens - when adequate tumor cellularity is present and decalcification has not been performed - are also more than adequate for additional molecular testing in lung cancer (12; 20; 21).

In conclusion, we have presented a five-year academic institutional experience relating the feasibility and success rates for lung cancer molecular testing of tissue obtained from a broad range of specimen types and modes of tissue acquisition. Importantly, since the molecular testing was not performed in-house, but rather by an independent commercial vendor, the results and conclusions drawn from this study are broadly applicable to other institutions or practices that also send out molecular testing of lung cancer specimens. Attention paid to specific specimen characteristics can help ensure optimal selection of tumor biopsies for successful genotyping.

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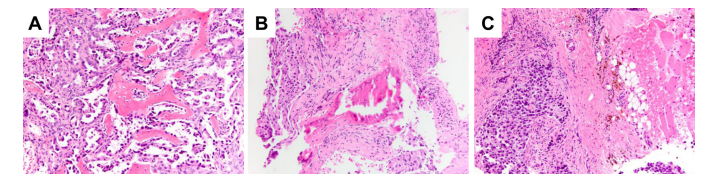


FIGURE 1.

Tumor specimens obtained from bony tissue. A: Failed genotyping of a core needle biopsy from a glenoid metastasis. Although the specimen demonstrates high tumor cellularity, the extensive infiltration of bone trabeculae necessitated decacification prior to processing. Subsequent genotyping from a second specimen (cell block from a transbronchial FNA) successfully demonstrated a deletion mutation in exon 19 of the EGFR gene. B: Successful tumor genotyping of a core needle biopsy from a L3 spinal metastasis in a 64 year old female never smoker. The poorly differentiated adenocarcinoma (5 and 9 o'clock) is seen in a background of fibrosis with only rare bony trabeculae, thus decalcification was unnecessary for specimen processing. Compare the appearance of the calcified bone trabeculae with the decalcified osteoid in part A; the former demonstrates shattering or fracturing artifact, and had a more basophilic hue. C: Successfully genotyped core needle biopsy specimen of a sacral metastasis. The poorly differentiated adenocarcinoma can be seen extending into the periosseous soft tissues (adipose tissue and skeletal muscle on the right). The lack of calcified tissue in this bony metastasis obviated the need for decalcification, leading to successful tumor genotyping. All images were taken at 200x original magnification, hematoxylin and eosin stain.

Table 1

Baseline characteristics of patients and tumors for which a sample was submitted for somatic genotyping technologies

	All patients(n=381)
Age at the time of biopsy mean(SD)	65.0(11.4)
Women n (%)	233(61.2)
Race n (%)	
White	289(75.9)
Asian	50(13.1)
Black	25(6.56)
Others	17(4.46)
Smoking status n (%)	
Current smoker	66(17.3)
Former smoker	209(54.9)
Never smoker	106(27.8)
Stage n (%)	<u> </u>
I	24(6.30)
II	22(5.77)
III	47(12.1)
IV	288(73.8)
Histology n (%)	
Adenocarcinoma	328(86.1)
Squamous cell carcinoma	11(2.89)
NSCLC (NOS)	39(10.2)
Others	3(0.78)
Anatomic site of biopsy n (%)	
Bone	23(6.04)
Brain	23(6.04)
Liver	14(3.67)
Lung	188(49.3)
Lymph node	85(22.3)
Pleura	45(11.8)
Others	3(0.79)
Type of biopsy n (%)	
Core needle biopsy	68(17.9)
Surgical	186(48.8)
Cell block from FNA	98(25.7)
Cell block from fluid	29(7.61)

	All patients(n=381)
EGFR mutation analysis Submitted samples n (%)	381(100)
KRAS mutation analysis Submitted samples n (%)	226(59.31)
ALK FISH analysis Submitted samples n (%)	275(72.17)

Table 2

Success and failure rates for EGFR mutation, KRAS mutation and ALK FISHtechniques in the submitted samples

All samples(n=381)			
	EGFR mutation	ALK FISH	KRAS mutation
Success n(%)*	359(94.2)	252(91.6)	207(91.6)
Positive/Mutated	86(22.6)	23(8.36)	71(31.4)
Negative/Wild-type	273(71.7)	229(83.3)	136(60.2)
Failure n(%)*	22(5.8)	23(8.36)	19(8.41)
Not done n(%)**	0(0)	106(27.8)	155(40.7)
Samples who underwe	nt all tests(n=207)		
	EGFR mutation	ALK FISH	KRAS mutation
Success n(%)*	191(92.3)	186(89.9)	190(91.8)
Positive/Mutated	32(15.5)	11(5.31)	65(31.4)
Negative/Wild-type	159(76.8)	175(84.5)	125(60.4)
Failure n(%)*	16(7.73)	21(10.1)	17(8.21)
Not done n(%)**	0(0)	0(0)	0(0)

^{*} Percentage out of the attempted assay

^{**}Percentage out of the number of all patients

Table 3

Characteristics of failure cases for EGFR mutation detection and their potential predictive factors by multivariable logistic regression

Predictive factors	No. of patients(%)	Multiv	ariable regre	ession
		Odds ratio	95% CI*	p value
Male	148(39)	1.00	0.57 to	0.45
Female	233(61)	1.42	3.58	
White	289(76)	1.21	0.39 to	0.74
Others	92(24)	1.00	3.64	
Non-smokers	106(28)	1.00	0.45 to	0.51
Smokers	275(72)	1.50	5.00	
Stage I-III	93(24)	1.11	0.29 to	0.88
Stage IV	288(76)	1.00	4.22	
Adenocarcinoma	328(86)	1.00	0.63 to	0.22
Others	53(14)	2.15	7.34	
Bone	23(6)	6.17	1.56 to	0.01
Others	358(94)	1.00	24.5	
Surgical	186(49)	1.00	3.58 to	0.002
Non-surgical	195(51)	28.5	226.2	

95%CI*: 95% Confidence interval.

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Table 4

Failure rates of each test stratified by the anatomic site sampled in the complete cohort

EGFR Mutation analysis (n=381)	ıtation aı	nalysis (n	=381)					
	Bone	Brain	Liver	Lung	Lymph node	Pleura	Others	Total
Failure n	4	0	2	11	2	2	1	22
%Total*	18.2	0	60.6	50.0	60.6	60.6	4.55	100.0
%Site**	17.4	0	14.3	5.85	2.35	4.44	33.3	5.77
Total	23	23	14	188	85	45	3	381
ALK FISH analysis (n=275)	analysis	(n=275)						
	Bone	Brain	Liver	Lung	Lymph node	Pleura	Others	Total
Failure n	4	0	1	12	4	1	1	23
"Total	17.4	0	4.35	52.2	17.4	4.35	4.35	100.0
%Site**	22.2	0	12.5	9.1	6.25	3.33	50.0	8.36
Total	18	21		132	64	30	2	275
KRAS mutation analysis (n=226)	ation ana	lysis (n=2	26)					
_	Bone	Brain	Liver	Lung	Lymph node	Pleura	Others	Total
Failure n	2	0	2	6	4	2	0	19
"Total	10.5	0	10.53	47.4	21.1	10.53	0.00	100.0
%Site**	14.3	0	50.0	8.11	7.41	8.00	0.00	8.41
Total	14	17	4	111	54	25	1	226

Recentage out of total success or failure

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^{**}Percentage out of the number of attempted biopsy on the each anatomic site

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Table 5

Failure rates of each test stratified by the anatomic site sampled in the core needle biopsy group

EGFK Mutation analysis (n=68)	itation a	nalysis (n	=68)				
	Bone	Liver	Lung (bronchos copy- guided)	Lung (image- guided percutaneous transthoracic)	Pleura	Others	Total
Failure n	4	2	2	7	0	1	16
%Total*	25.0	12.5	12.5	43.8	0	6.25	100.0
%Site**	4.4.4	14.3	11.1	31.8	0	50.0	23.5
Total	6_	14	18	22	3	2	89_
ALK FISH	analysis (n=53)	(n=53)					
	Bone	Liver	Lung (bronchos Copy- guided)	Lung (image- guided percutaneous transthoracic)	Pleura	Others	Total
Failure n	4	1	2	9	0	1	14
"Total"	28.6	7.14	14.3	42.9	0	7.14	100.0
%Site**	44.4	12.5	12.5	35.3	0	100	26.4
Total	6	_8_	16	17	2	1	53
KRAS mutation analysis (n=41)	ation ana	lysis (n=4	(1)				
	Bone	Liver	Lung (bronchos Copy- guided)	Lung (image- guided percutaneous transthoracic)	Pleura	Others	Total
Failure n	2	2	1	3	0	1	6
%Total*	22.2	22.2	11.1	33.3	0	11.1	100.0
%Site**	25.0	50.0	29.9	27.3	0	100	22.0
Total	~	4	15	11	2	1	41

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Percentage out of the number of attempted biopsy on the each anatomic site

* Percentage out of total success or failure