

# Template for Reporting Results of Biomarker Testing of Specimens From Patients With Non-Small Cell Carcinoma of the Lung

Version: 2.0.1.0

Protocol Posting Date: November 2021

This biomarker template is not required for accreditation purposes but may be used to facilitate compliance with CAP Accreditation Program Requirements

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With guidance from the CAP Cancer and CAP Pathology Electronic Reporting Committees.

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#### **Accreditation Requirements**

Completion of the template is the responsibility of the laboratory performing the biomarker testing and/or providing the interpretation. When both testing and interpretation are performed elsewhere (eg, a reference laboratory), synoptic reporting of the results by the laboratory submitting the tissue for testing is also encouraged to ensure that all information is included in the patient's medical record and thus readily available to the treating clinical team. This template is not required for accreditation purposes.

#### **Summary of Changes**

## v 2.0.1.0

Changed Questions for RET and BRAF Interpretation from select all that apply to single select

# **Reporting Template**

**Protocol Posting Date: November 2021** 

Select a single response unless otherwise indicated.

# **CASE SUMMARY: (Lung Biomarker Reporting Template)**

Completion of the template is the responsibility of the laboratory performing the biomarker testing and / or providing the interpretation. When both testing and interpretation are performed elsewhere (e.g., a reference laboratory), synoptic reporting of the results by the laboratory submitting the tissue for testing is also encouraged to ensure that all information is included in the patient's medical record and thus readily available to the treating clinical team.

Gene names should follow recommendations of The Human Genome Organisation (HUGO) Nomenclature Committee (www.genenames.org; accessed February 10, 2015).

All reported gene sequence variations should be identified following the recommendations of the Human Genome Variation Society (www.hgvs.org/mutnomen/; accessed February 10, 2015).

### **SPECIMEN**

+Adequacy of Sample for Testing
Adequate +Estimated % Tumor Cellularity (area used for testing): %
Suboptimal (explain):
Gabapamar (explain).
+Specimen Type
Untreated diagnostic specimen
Relapse specimen (after treatment; specify)#:
# When data is available, specify treatment type. This is most relevant to targeted inhibitors associated with specific genomic changes conferring treatment resistance.
RESULTS EGFR
+Mutational Analysis
No EGFR mutation detected
Mutation(s) identified
EGFR:p.G719X
EGFR Exon 19 deletion (specify if known):
EGFR Exon 20 insertion (specify if known):
EGFR:p.S768I
EGFR:p.T790M
EGFR:p.L858R
EGFR:p.L861Q
Other (specify):
Cannot be determined (explain):
+EGFR L858R by Immunohistochemistry (clone 43B2)
Negative
Positive
Equivocal (explain):
+EGFR Exon 19 Deletion (E746_A750del) (clone 6B6)
Negative
Positive

Equivocal (explain):
+Interpretation (select all that apply)
An EGFR mutation is present that is associated with response to EGFR tyrosine kinase inhibitors
An EGFR mutation is present that is associated with resistance to EGFR tyrosine kinase inhibitors
Two EGFR mutations are present, one of which is associated with resistance to EGFR tyrosine
kinase inhibitors
EGFR L858R immunohistochemical staining is positive, which is associated with response to
EGFR tyrosine kinase inhibitors
EGFR E746_A750del immunohistochemical staining is positive, which is associated with response
to EGFR tyrosine kinase inhibitors
ALK
+Rearrangement by Molecular Methods
No ALK rearrangement detected
Rearrangement identified
EML4-ALK (specify variant type, if known):
KIF5B-ALK
KLC1-ALK
Other ALK rearrangement (specify if known): :
Cannot be determined (explain):
+ALK Immunohistochemistry
Negative
Positive
Equivocal (explain):
that are not affine to all and all the store all the
+Interpretation (select all that apply)
An ALK fusion is identified that is associated with response to ALK tyrosine kinase inhibitors
ALK immunohistochemical staining is positive which is associated with response to ALK tyrosine
kinase inhibitors
ROS1
+Rearrangement by Molecular Methods
No ROS1 rearrangement detected
ROS1 rearrangement identified
Cannot be determined (explain):
carnot be determined (explain).
+ROS1 by Immunohistochemistry
Negative
Positive
Equivocal (explain):
+Interpretation (select all that apply)
A ROS1 fusion is present, which is associated with response to ROS tyrosine kinase inhibitors
ROS1 immunohistochemical staining is positive, which is associated with response to ROS1
tyrosine kinase inhibitors
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+Rearrangement by Molecular Methods
No RET rearrangement detected
RET rearrangement identified
Cannot be determined (explain):
+Interpretation
A RET fusion is present which is associated with response to RET tyrosine kinase inhibitors
No RET fusions are detected
KRAS
+Mutational Analysis
No KRAS mutation detected
Mutation(s) identified
KRAS:p.G12C
KRAS:p.G12D
KRAS:p.G12V
KRAS:p.G12S
KRAS:p.G12A
KRAS:p.G12R
KRAS:p.G13D
KRAS:p.G13C
KRAS:p.Q61L
Other (specify):
Cannot be determined (explain):
<ul> <li>+Interpretation (select all that apply)</li> <li> A KRAS mutation is identified which is associated with resistance to tyrosine kinase inhibitor therapy</li> <li> A KRAS mutation is identified which is associated with response to specific inhibitors</li> </ul>
BRAF
+Mutational Analysis
No BRAF mutations detected
Mutation(s) identified
BRAF:p.V600E
Other (specify):
Cannot be determined (explain):
+Interpretation
A BRAF mutation is present which is associated with response to BRAF inhibitors
No BRAF mutations are detected
ERBB2
+Mutational Analysis
No ERBB2 mutations detected
Mutation(s) identified
ERBB2:p.S310F
ERBB2:p.L755S

ERBB2:p.Y772_A775dup insertion Other (specify):	
Cannot be determined (explain):	_
+Copy Number Analysis  No ERBB2 (HER2) amplification detected  ERBB2 (HER2) amplification identified	
Specify Copy Number: Specify Ratio to Centromere 17: Cannot be determined (explain):	
+HER2 immunohistochemistry	_
Negative (0-1)	
Equivocal (2+)	
Positive (3+) Cannot be determined (explain):	_
<ul> <li>+Interpretation (select all that apply)</li> <li>An ERBB2 (HER2) mutation is present which is assometed.</li> <li>ERBB2 (HER2) amplification is present which is assometed.</li> <li>HER2 is positive by immunohistochemistry (3+) which therapy.</li> </ul>	ciated response to anti-HER2 therapy
MET	
+Mutational Analysis  No MET mutation detected	
Mutation(s) identified	
MET:p.D963_splice mutation	
MET:p.D1010N MET:p.D1010_splice mutation	
MET exon 14 deletion	
Other (specify):	
Cannot be determined (explain):	_
+Copy Number Analysis  No MET amplification detected	
MET amplification identified	
Specify Copy Number:	
Specify Ratio to Centromere 7:	
Cannot be determined (explain):	_
+Interpretation (select all that apply)	
A MET alteration is present which is associated with MET amplification is present which is associated with	
NTRK	
+Rearrangement by Molecular Methods	
No NTRK rearrangement detected NTRK rearrangement identified (specify if known):	
INTER TEATRAINCEITIENT (QUEITUILEU (SDECIIV II KNOWN).	

Cannot be determined (explain):
+NTRK by immunohistochemistry
Negative
Positive
Equivocal
+Interpretation (select all that apply)
An NTRK fusion is present which is associated with response to NTRK inhibitors
NTRK immunohistochemical staining is present. Fusion testing by NGS or FISH will be performed
NTRK immunohistochemical staining is present but fusion testing is negative. This is not associated with response to NTRK inhibitors
associated with response to NTAX inhibitors
Mismatch Repair
+Immunohistochemistry (IHC) Testing for Mismatch Repair (MMR) Proteins (select all that
apply) MLH1
MLH1 Result
Intact nuclear expression
Loss of nuclear expression
Cannot be determined (explain):
MSH2
MSH2 Result
Intact nuclear expression
Loss of nuclear expression
Cannot be determined (explain):
MSH6
MSH6 Result
Intact nuclear expression
Loss of nuclear expression
Cannot be determined (explain):
PMS2
PMS2 Result
Intact nuclear expression
Loss of nuclear expression
Cannot be determined (explain):
Background nonneoplastic tissue / internal control with intact nuclear expression
+Microsatellite Instability (MSI)
MSI-Stable (MSS)
MSI-Low (MSI-L)
MSI-High (MSI-H)
Cannot be determined:

+Interpretation (select all that apply)	
The case is MSI-H which is associated with response to immune checkpoint inhibitors	
The case is mismatch repair deficient which is associated with response to immune checkpoin	ıt
inhibitors	
Tumor Mutational Burden	
+Specify Tumor Mutational Burden:	
+Tumor Mutational Burden Level	
Low	
High	
Equivocal	
Cannot be determined (explain):	
+Interpretation	
The case is TMB-high which is associated with response to immune checkpoint inhibitors	
The case is TMB low; this finding is not associated with response to immune checkpoint inhibition.	tors
PD-L1 IHC	
+PD-L1 IHC Interpretation	
Positive	
Negative	
Cannot be determined (indeterminate)	
+Specify Percentage of Tumor Cells with Staining (TPS): %	
+Combined Number of Tumor and Immune Cells with Staining per 100 Tumor Cells (CPS):	
+Specify Percentage of Tumor-associated Immune Cells with Staining:	%
+Specify Percentage of Tumor Area Occupied by Tumor-associated Immune Cells:	
+Comments:	
- Comments.	
Methods	
+Antibody	
22C3	
SP142	
SP263	
28-8	
Other (specify):	
+Controls (select all that apply)	
Internal control cells present; expected immunoreactivity	
Internal control cells present; no immunoreactivity of either tumor cells or internal controls	
External controls available, expected immunoreactivity	
External controls available; no immunoreactivity in expected cells	

+Assay Information Food and Drug Administration (FDA) cleared test / vendor (specify): Laboratory-developed test	
+Specify Quantitative Imaging Analytics Performed:	
Other Markers Tested (repeat as needed) +Specify Other Marker and Results:	
COMMENTS	
Comment(s):	