

Department of Pathology and Laboratory Medicine No.201, Sec. 2, Shipai Rd., Beitou District, Taipei City, Taiwan 11217, R.O.C. Tel: 02-2875-7449

Date: 11 Sep 2020

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Sample Information

Patient Name: 林義賢 Gender: Male ID No.: Q120542816 History No.: 46460462

Age: 58

Ordering Doctor: DOC3153J 黄煦晴

Ordering REQ.: 0AVUETR Signing in Date: 2020/09/09

Path No.: S109-99982 **MP No.:** F20070

Assay: Oncomine Focus Assay

Sample Type: FFPE Block No.: \$109-28573A Percentage of tumor cells: 90%

Note:

Sample Cancer Type: Non-Small Cell Lung Cancer

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Report Highlights

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Relevant Non-Small Cell Lung Cancer Findings

Gene	Finding	Gene	Finding
ALK	Not detected	NTRK1	Not detected
BRAF	Not detected	NTRK2	Not detected
EGFR	Not detected	NTRK3	Not detected
ERBB2	Not detected	RET	Not detected
KRAS	KRAS p.(Q61L) c.182_183delAAinsTT	ROS1	Not detected
MET	Not detected		



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Relevant Biomarkers

Tier	Genomic Alteration	Relevant Therapies (In this cancer type)	Relevant Therapies (In other cancer type)	Clinical Trials
IA	KRAS p.(Q61L) c.182_183delAAinsTT	None	cabozantinib	32
	KRAS proto-oncogene, GTPase Allele Frequency: 53.81%			

Public data sources included in relevant therapies: FDA1, NCCN, EMA2, ESMO

Tier Reference: Li et al. Standards and Guidelines for the Interpretation and Reporting of Sequence Variants in Cancer: A Joint Consensus Recommendation of the Association for Molecular Pathology, American Society of Clinical Oncology, and College of American Pathologists. J Mol Diagn. 2017 Jan;19(1):4-23.

Variant Details

DNA Sequence Variants								
Gene	Amino Acid Change	Coding	Variant ID	Locus	Allele Frequency	Transcript	Variant Effect	Coverage
KRAS	p.(Q61L)	c.182_183delAAinsT T		chr12:25380275	53.81%	NM_033360.3	missense	1994
JAK1	p.(=)	c.2199A>G		chr1:65310489	37.34%	NM_002227.3	synonymous	1995
ALK	p.(D1529E)	c.4587C>G		chr2:29416366	61.43%	NM_004304.4	missense	1999
ALK	p.(I1461V)	c.4381A>G		chr2:29416572	99.95%	NM_004304.4	missense	1998
ALK	p.(=)	c.3600G>C		chr2:29443617	59.01%	NM_004304.4	synonymous	1998
ALK	p.(=)	c.3375C>A		chr2:29445458	60.12%	NM_004304.4	synonymous	1996
FGFR3	p.(=)	c.1953G>A		chr4:1807894	99.70%	NM_000142.4	synonymous	1998
PDGFRA	p.(=)	c.939T>G		chr4:55133726	45.38%	NM_006206.5	synonymous	1992
PDGFRA	p.(=)	c.1701A>G		chr4:55141055	99.80%	NM_006206.5	synonymous	2000
PDGFRA	p.(=)	c.2472C>T		chr4:55152040	46.55%	NM_006206.5	synonymous	2000
KIT	p.(=)	c.1638A>G		chr4:55593481	46.47%	NM_000222.2	synonymous	1999
FGFR4	p.(P136L)	c.407C>T		chr5:176517797	99.10%	NM_213647.2	missense	1999
EGFR	p.(=)	c.2361G>A		chr7:55249063	40.40%	NM_005228.4	synonymous	2000
RET	p.(=)	c.2307G>T		chr10:43613843	77.37%	NM_020975.4	synonymous	1997
RET	p.(=)	c.2712C>G		chr10:43615633	77.42%	NM_020975.4	synonymous	1997

Biomarker Descriptions

KRAS (KRAS proto-oncogene, GTPase)

Background: The KRAS proto-oncogene encodes a GTPase that functions in signal transduction and is a member of the RAS superfamily which also includes NRAS and HRAS. RAS proteins mediate the transmission of growth signals from the cell surface to the nucleus via the PI3K/AKT/MTOR and RAS/RAF/MEK/ERK pathways, which regulate cell division, differentiation, and survival^{1,2,3}.



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A Both for use and

contraindicated

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No evidence

(I/II)

×

×

Biomarker Descriptions (continued)

Alterations and prevalence: Recurrent mutations in RAS oncogenes cause constitutive activation and are found in 20-30% of cancers. KRAS mutations are observed in up to 10-20% of uterine cancer, 30-35% of lung adenocarcinoma and colorectal cancer, and about 60% of pancreatic cancer⁴. The majority of KRAS mutations consist of point mutations occurring at G12, G13, and Q61^{4,5,6}. Mutations at A59, K117, and A146 have also been observed but are less frequent^{7,8}.

Potential relevance: Currently, no therapies are approved for KRAS aberrations. However, the KRAS G12C inhibitor, sotorasib (AMG $\overline{510}$) 9 , was granted fast track designation (2019) for previously treated non-small cell lung cancer (NSCLC) patients with KRAS G12C mutations. The EGFR antagonists, cetuximab 10 and panitumumab 11 , are contraindicated for treatment of colorectal cancer patients with KRAS mutations in exon 2 (codons 12 and 13), exon 3 (codons 59 and 61), and exon 4 (codons 117 and 146) 8 . Additionally, KRAS mutations are associated with poor prognosis in NSCLC 12 .

In this cancer type and

other cancer types

Contraindicated

Relevant Therapy Summary

In this cancer type In other cancer

afatinib + selumetinib

type

Relevant Therapy	FDA	NCCN	EMA	ESMO	Clinical Trials*
cetuximab	0	0	0	0	×
panitumumab	0	0	×	0	×
cetuximab + oxaliplatin	×	×	0	×	×
panitumumab + oxaliplatin	×	×	0	×	×
cabozantinib	×	×	×	0	×
cetuximab + chemotherapy	×	×	×	0	×
panitumumab + chemotherapy	×	×	×	0	×
bevacizumab, chemotherapy	×	×	×	×	(III)
lenvatinib, pembrolizumab, chemotherapy	×	×	×	×	(III)
atezolizumab, cobimetinib	×	×	×	×	(II)
regorafenib, chemotherapy	×	×	×	×	(II)
selumetinib, ulixertinib	×	×	×	×	(II)
sintilimab, anlotinib hydrochloride	×	×	×	×	(II)
spartalizumab	×	×	×	×	(II)
targeted therapy, chemotherapy	×	×	×	×	(II)
TVB-2640	×	×	×	×	(II)

* Most advanced phase (IV, III, II/III, II, I/II, I) is shown and multiple clinical trials may be available.

×

×

Disclaimer: The data presented here is from a curated knowledgebase of publicly available information, but may not be exhaustive. The data version is 2020.06(005).

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Relevant Therapy Summary (continued)

In this cancer type O In other cancer

type

In this cancer type and other cancer types

Contraindicated

A Both for use and contraindicated

X No evidence

KRAS p.(Q61L) c.182_183delAAinsTT (continued)

Relevant Therapy	FDA	NCCN	EMA	ESMO	Clinical Trials*
ASTX029	×	×	×	×	(I/II)
avelumab, binimetinib, talazoparib	×	×	×	×	(I/II)
binimetinib + palbociclib, binimetinib, palbociclib	×	×	×	×	(I/II)
lapatinib, trametinib	×	×	×	×	(/)
mirdametinib, lifirafenib	×	×	×	×	(/)
navitoclax, trametinib	×	×	×	×	(/)
neratinib, valproic acid	×	×	×	×	(/)
RMC-4630, cobimetinib	×	×	×	×	(/)
selinexor, chemotherapy	×	×	×	×	(/)
selumetinib, durvalumab, tremelimumab	×	×	×	×	(I/II)
zotatifin	×	×	×	×	(/)
BGB-3245	×	×	×	×	(I)
cobimetinib, belvarafenib	×	×	×	×	(I)
LXH254 , LTT-462, trametinib, ribociclib	×	×	×	×	(I)
LXH254 , spartalizumab	×	×	×	×	(I)
LY3214996, midazolam, abemaciclib, chemotherapy, encorafenib, cetuximab	×	×	×	×	(1)
NBF-006	×	×	×	×	(I)
neratinib, trametinib	×	×	×	×	(I)
pembrolizumab + trametinib	×	×	×	×	(I)
RMC-4630	×	×	×	×	(I)
RO-5126766	×	×	×	×	(I)
TAK 659, chemotherapy	×	×	×	×	(l)

^{*} Most advanced phase (IV, III, II/II, II, I/II, I) is shown and multiple clinical trials may be available.



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Relevant Therapy Details

Current FDA Information

In this cancer type	O In other cancer type	0	In this cancer type and	(Ontraindicated	Not recommended	D	Resistance
,,	,,		other cancer types					

FDA information is current as of 2020-05-26. For the most up-to-date information, search www.fda.gov.

KRAS p.(Q61L) c.182_183delAAinsTT

cetuximab

Cancer type: Colorectal Cancer Label as of: 2019-04-23 Variant class: KRAS Q61 mutation

Indications and usage:

Erbitux® is an epidermal growth factor receptor (EGFR) antagonist indicated for treatment of:

Head and Neck Cancer

- Locally or regionally advanced squamous cell carcinoma of the head and neck in combination with radiation therapy.
- Recurrent locoregional disease or metastatic squamous cell carcinoma of the head and neck in combination with platinumbased therapy with fluorouracil.
- Recurrent or metastatic squamous cell carcinoma of the head and neck progressing after platinum-based therapy.

Colorectal Cancer

K-Ras wild-type, EGFR-expressing, metastatic colorectal cancer as determined by FDA-approved test

- in combination with FOLFIRI for first-line treatment,
- in combination with irinotecan in patients who are refractory to irinotecan-based chemotherapy,
- as a single agent in patients who have failed oxaliplatin- and irinotecan-based chemotherapy or who are intolerant to irinotecan.

Limitations of Use: Erbitux® is not indicated for treatment of Ras-mutant colorectal cancer or when the results of the Ras mutation tests are unknown.

Reference:

https://www.accessdata.fda.gov/drugsatfda_docs/label/2019/125084s273lbl.pdf



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KRAS p.(Q61L) c.182_183delAAinsTT (continued)

panitumumab

Cancer type: Colorectal Cancer Label as of: 2017-06-29 Variant class: KRAS Q61 mutation

Indications and usage:

VECTIBIX® is an epidermal growth factor receptor (EGFR) antagonist indicated for the treatment of wild-type RAS (defined as wild-type in both KRAS and NRAS as determined by an FDA-approved test for this use) metastatic colorectal cancer (mCRC):

- In combination with FOLFOX for first-line treatment.
- As monotherapy following disease progression after prior treatment with fluoropyrimidine, oxaliplatin, and irinotecancontaining chemotherapy.
- Limitation of Use: VECTIBIX® is not indicated for the treatment of patients with RAS-mutant mCRC or for whom RAS mutation status is unknown.

Reference:

https://www.accessdata.fda.gov/drugsatfda_docs/label/2017/125147s207lbl.pdf



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Current NCCN Information

In this cancer type O In other cancer type

In this cancer type and other cancer types

Contraindicated

Not recommended Resistance

NCCN information is current as of 2020-05-01. For the most up-to-date information, search www.nccn.org. For NCCN International Adaptations & Translations, search www.nccn.org/global/international_adaptations.aspx.

KRAS p.(Q61L) c.182_183delAAinsTT

cetuximab

Cancer type: Colon Cancer Variant class: KRAS exon 3 mutation

Summary:

NCCN Guidelines® include the following supporting statement(s):

"Patients with any known KRAS mutation (exon 2, 3, 4) or NRAS mutation (exon 2, 3, 4) should not be treated with either cetuximab or panitumumab."

Reference: NCCN Guidelines® - NCCN-Colon Cancer [Version 3.2020]

cetuximab

Variant class: KRAS exon 3 mutation Cancer type: Rectal Cancer

Summary:

NCCN Guidelines® include the following supporting statement(s):

"Patients with any known KRAS mutation (exon 2, 3, 4) or NRAS mutation (exon 2, 3, 4) should not be treated with either cetuximab or panitumumab."

Reference: NCCN Guidelines® - NCCN-Rectal Cancer [Version 4.2020]

panitumumab

Variant class: KRAS exon 3 mutation Cancer type: Colon Cancer

Summary:

NCCN Guidelines® include the following supporting statement(s):

"Patients with any known KRAS mutation (exon 2, 3, 4) or NRAS mutation (exon 2, 3, 4) should not be treated with either cetuximab or panitumumab."

Reference: NCCN Guidelines® - NCCN-Colon Cancer [Version 3.2020]

panitumumab

Variant class: KRAS exon 3 mutation Cancer type: Rectal Cancer

Summary:

NCCN Guidelines® include the following supporting statement(s):

"Patients with any known KRAS mutation (exon 2, 3, 4) or NRAS mutation (exon 2, 3, 4) should not be treated with either cetuximab or panitumumab."

Reference: NCCN Guidelines® - NCCN-Rectal Cancer [Version 4.2020]



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KRAS p.(Q61L) c.182_183delAAinsTT (continued)

EGFR tyrosine kinase inhibitor

Cancer type: Non-Small Cell Lung Cancer Variant class: KRAS mutation

Summary:

NCCN Guidelines® include the following supporting statement(s):

- "Non-responsiveness to EGFR TKI therapy is associated with KRAS and BRAF mutations and ALK or ROS1 gene fusions."
- "KRAS mutational status is also predictive of lack of therapeutic efficacy with EGFR TKIs."

Reference: NCCN Guidelines® - NCCN-Non-Small Cell Lung Cancer [Version 4.2020]



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Current EMA Information

EMA information is current as of 2020-05-26. For the most up-to-date information, search www.ema.europa.eu/ema.

KRAS p.(Q61L) c.182_183delAAinsTT

cetuximab, cetuximab + oxaliplatin

Cancer type: Colorectal Cancer Label as of: 2020-01-30 Variant class: KRAS exon 3 mutation

Reference:

https://www.ema.europa.eu/en/documents/product-information/erbitux-epar-product-information_en.pdf

panitumumab + oxaliplatin

Cancer type: Colorectal Cancer Label as of: 2020-01-24 Variant class: KRAS exon 3 mutation

Reference:

https://www.ema.europa.eu/en/documents/product-information/vectibix-epar-product-information_en.pdf



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Current ESMO Information

In this cancer type O In other cancer type

In this cancer type and other cancer types

Contraindicated

Not recommended Resistance

ESMO information is current as of 2020-05-01. For the most up-to-date information, search www.esmo.org.

KRAS p.(Q61L) c.182_183delAAinsTT

O cabozantinib

Cancer type: Thyroid Gland Medullary Carcinoma Variant class: RAS mutation

ESMO Level of Evidence/Grade of Recommendation: II / C

Population segment (Line of therapy):

Metastatic Thyroid Gland Medullary Carcinoma (First-line therapy)

Reference: ESMO Clinical Practice Guidelines - ESMO-Thyroid Cancer [Annals of Oncology 30: 1856-1883, 2019 doi:10.1093/ annonc/mdz4001

cetuximab

Cancer type: Colorectal Cancer Variant class: KRAS exon 3 mutation

Summary:

ESMO Clinical Practice Guidelines include the following supporting statement:

"It has been demonstrated that the (potential) benefit of anti-EGFR antibodies in all treatment lines and either as a single agent or in combination with any chemotherapy regimen is limited to patients in whom a RAS mutation is excluded. It was shown that the 'expanded RAS' analysis (also including the detection of mutations in exons 3 and 4 of the KRAS gene as well as mutations in the NRAS [exons 2-4] gene) is superior to the KRAS (exon 2) analysis in predicting both more efficacy in the expanded RAS wild-type (WT) patients and a potential detrimental effect in patients harbouring any RAS mutation in their tumour genome [II/A].'

Reference: ESMO Clinical Practice Guidelines - ESMO-Metastatic Colorectal Cancer [Ann Oncol (2014) 25 (suppl 3): iii1-iii9. (eUpdate: 20 September 2016; Corrigendum: 21 July 2015)]



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KRAS p.(Q61L) c.182_183delAAinsTT (continued)

cetuximab + chemotherapy

Cancer type: Colorectal Cancer Variant class: KRAS exon 3 mutation

Summary:

ESMO Clinical Practice Guidelines include the following supporting statement:

- "It has been demonstrated that the (potential) benefit of anti-EGFR antibodies in all treatment lines and either as a single agent or in combination with any chemotherapy regimen is limited to patients in whom a RAS mutation is excluded. It was shown that the 'expanded RAS' analysis (also including the detection of mutations in exons 3 and 4 of the KRAS gene as well as mutations in the NRAS [exons 2-4] gene) is superior to the KRAS (exon 2) analysis in predicting both more efficacy in the expanded RAS wild-type (WT) patients and a potential detrimental effect in patients harbouring any RAS mutation in their tumour genome [II/A]."
- "Thus, the activity of the anti-EGFR antibodies is confined to RAS WT tumours (and not only KRAS WT tumours). This is true for the combinations of cetuximab or panitumumab alone or with irinotecan- and oxaliplatin-based regimens. Treatment with anti-EGFR antibodies may even harm patients with a RAS mutation, especially when combined with oxaliplatin [I/A]."

Reference: ESMO Clinical Practice Guidelines - ESMO-Metastatic Colorectal Cancer [Ann Oncol (2014) 25 (suppl 3): iii1-iii9. (eUpdate: 20 September 2016; Corrigendum: 21 July 2015)]

panitumumab

Cancer type: Colorectal Cancer Variant class: KRAS exon 3 mutation

Summary:

ESMO Clinical Practice Guidelines include the following supporting statement:

■ "It has been demonstrated that the (potential) benefit of anti-EGFR antibodies in all treatment lines and either as a single agent or in combination with any chemotherapy regimen is limited to patients in whom a RAS mutation is excluded. It was shown that the 'expanded RAS' analysis (also including the detection of mutations in exons 3 and 4 of the KRAS gene as well as mutations in the NRAS [exons 2-4] gene) is superior to the KRAS (exon 2) analysis in predicting both more efficacy in the expanded RAS wild-type (WT) patients and a potential detrimental effect in patients harbouring any RAS mutation in their tumour genome [II/A]."

Reference: ESMO Clinical Practice Guidelines - ESMO-Metastatic Colorectal Cancer [Ann Oncol (2014) 25 (suppl 3): iii1-iii9. (eUpdate: 20 September 2016; Corrigendum: 21 July 2015)]



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KRAS p.(Q61L) c.182_183delAAinsTT (continued)

panitumumab + chemotherapy

Cancer type: Colorectal Cancer Variant class: KRAS exon 3 mutation

Summary:

ESMO Clinical Practice Guidelines include the following supporting statement:

- "It has been demonstrated that the (potential) benefit of anti-EGFR antibodies in all treatment lines and either as a single agent or in combination with any chemotherapy regimen is limited to patients in whom a RAS mutation is excluded. It was shown that the 'expanded RAS' analysis (also including the detection of mutations in exons 3 and 4 of the KRAS gene as well as mutations in the NRAS [exons 2-4] gene) is superior to the KRAS (exon 2) analysis in predicting both more efficacy in the expanded RAS wild-type (WT) patients and a potential detrimental effect in patients harbouring any RAS mutation in their tumour genome [II/A]."
- "Thus, the activity of the anti-EGFR antibodies is confined to RAS WT tumours (and not only KRAS WT tumours). This is true for the combinations of cetuximab or panitumumab alone or with irinotecan- and oxaliplatin-based regimens. Treatment with anti-EGFR antibodies may even harm patients with a RAS mutation, especially when combined with oxaliplatin [I/A]."

Reference: ESMO Clinical Practice Guidelines - ESMO-Metastatic Colorectal Cancer [Ann Oncol (2014) 25 (suppl 3): iii1-iii9. (eUpdate: 20 September 2016; Corrigendum: 21 July 2015)]



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Signatures
Testing Personnel:
Laboratory Supervisor:
Pathologist:



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