

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

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Sample Information

Patient Name: 葉銀清 Gender: Male ID No.: F101723571 History No.: 12442614

Age: 64

Ordering Doctor: DOC3064F 陳育民

Ordering REQ.: D5DC39F Signing in Date: 2020/09/24

Path No.: \$109-89666 **MP No.:** F20076

Assay: Oncomine Focus Assay

Sample Type: FFPE Block No.: \$109-77933A Percentage of tumor cells: 50%

Note:

Sample Cancer Type: Non-Small Cell Lung Cancer

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| BioBank with >1% allele frequency) | |
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Report Highlights

1 Relevant Biomarkers1 Therapies Available35 Clinical Trials

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.(G12V) c.35G>T | 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.(G12V) c.35G>T | None | cabozantinib | 35 |
| | KRAS proto-oncogene, GTPase | | | |
| | Allele Frequency: 24.80% | | | |

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.

Variants (Exclude variant in Taiwan BioBank with >1% allele frequency)

| DNA | Sequence Varia | ants | | | | | | |
|------|-------------------|---------|------------|----------------|---------------------|-------------|----------------|----------|
| Gene | Amino Acid Change | Coding | Variant ID | Locus | Allele Frequency | Transcript | Variant Effect | Coverage |
| KRAS | p.(G12V) | c.35G>T | COSM520 | chr12:25398284 | 24.80% | NM_033360.3 | missense | 1992 |

Biomarker Descriptions

KRAS (KRAS proto-oncogene, GTPase)

<u>Background:</u> 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}.

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 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, cetuximab¹⁰ and panitumumab¹¹, 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 NSCLC¹².

Relevant Therapy Summary

| In this cancer type O In oth type | her cancer In this cancer ty other cancer typ | pe and O Contrain | | for use and Xaindicated | No evidence |
|-----------------------------------|--------------------------------------------------|-------------------|--------|-------------------------|------------------|
| KRAS p.(G12V) c.35 | G>T | | | | |
| Relevant Therapy | | FDA NC | CN EMA | ESMO | Clinical Trials* |
| cetuximab | | 0 0 | 0 | 0 | × |

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



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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.(G12V) c.35G>T (continued)

| Relevant Therapy | FDA | NCCN | EMA | ESMO | Clinical Trials* |
|-----------------------------------------------------|-----|------|-----|------|------------------|
| 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) |
| afatinib + selumetinib | × | × | × | × | (I/II) |
| anti-KRAS G12V mTCR | × | × | × | × | (/) |
| ASTX029 | × | × | × | × | (/) |
| avelumab, binimetinib, talazoparib | × | × | × | × | (/) |
| binimetinib + palbociclib, binimetinib, palbociclib | × | × | × | × | (1/11) |
| lapatinib, trametinib | × | × | × | × | (/) |
| mirdametinib, lifirafenib | × | × | × | × | (1/11) |
| navitoclax, trametinib | × | × | × | × | (/) |
| neratinib, valproic acid | × | × | × | × | (1/11) |
| RMC-4630, cobimetinib | × | × | × | × | (I/II) |

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

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(I)

(I)

×

×



RO-5126766

TAK 659, chemotherapy

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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

×

×

×

×

Both for use and contraindicated

X No evidence

| Relevant Therapy | FDA | NCCN | EMA | ESMO | Clinical Trials* |
|-------------------------------------------------------------------------|-----|------|-----|------|------------------|
| selinexor, chemotherapy | × | × | X | × × | |
| · · | | | | | |
| selumetinib, durvalumab, tremelimumab | × | × | × | × | (1/11) |
| zotatifin | × | × | × | × | (1/11) |
| BGB-3245 | × | × | × | × | (l) |
| cobimetinib, belvarafenib | × | × | × | × | (I) |
| JAB-3312 | × | × | × | × | (I) |
| LXH254 , LTT-462, trametinib, ribociclib | × | × | × | × | (I) |
| LXH254 , spartalizumab | × | × | × | × | (I) |
| LY3214996, midazolam, abemaciclib, chemotherapy, encorafenib, cetuximab | × | × | × | × | ● (l) |
| mRNA-5671, pembrolizumab | × | × | × | × | (I) |
| NBF-006 | × | × | × | × | (I) |
| neratinib, trametinib | × | × | × | × | (I) |
| pembrolizumab + trametinib | × | × | × | × | (I) |
| RMC-4630 | × | × | × | × | (I) |

×

×

^{*} 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 and other cancer types

Ontraindicated

Not recommended

Resistance

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

KRAS p.(G12V) c.35G>T

cetuximab

Cancer type: Colorectal Cancer Label as of: 2019-04-23 Variant class: KRAS G12 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.(G12V) c.35G>T (continued)

panitumumab

Cancer type: Colorectal Cancer Label as of: 2017-06-29 Variant class: KRAS G12 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.(G12V) c.35G>T

cetuximab

Cancer type: Colon Cancer Variant class: KRAS exon 2 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 2 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 2 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 2 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.(G12V) c.35G>T (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

In this cancer type and O Contraindicated other cancer types

Not recommended Resistance

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

KRAS p.(G12V) c.35G>T

cetuximab, cetuximab + oxaliplatin

Cancer type: Colorectal Cancer Label as of: 2020-01-30 Variant class: KRAS exon 2 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 2 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.(G12V) c.35G>T

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 2 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.(G12V) c.35G>T (continued)

cetuximab + chemotherapy

Cancer type: Colorectal Cancer Variant class: KRAS exon 2 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 turnour 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 2 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.(G12V) c.35G>T (continued)

panitumumab + chemotherapy

Cancer type: Colorectal Cancer Variant class: KRAS exon 2 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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