Taipei Veterans General Hospital



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: 15 Oct 2020

1 of 8

Sample Information

Patient Name: 劉秉成

Gender: Male **ID No.:** A100808105 **History No.:** 796612

Age: 78

Ordering Doctor: DOC3064F 陳育民

Ordering REQ.: C21PL4K Signing in Date: 2020/10/14

Path No.: \$109-89734 **MP No.:** F20085

Assay: Oncomine Focus Assay

Sample Type: FFPE Block No.: \$109-32378A Percentage of tumor cells: 50%

Note:

Sample Cancer Type: Non-Small Cell Lung Cancer

Table of Contents	Page
Variant Details	2
Biomarker Descriptions	2
Relevant Therapy Summary	3
Relevant Therapy Details	5

Report Highlights

1 Relevant Biomarkers1 Therapies Available26 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	ERBB2 exon 20 insertion	RET	Not detected	
KRAS	Not detected	ROS1	Not detected	
MET	Not detected			



Tel: 02-2875-7449

Date: 15 Oct 2020 2 of 8

Relevant Biomarkers

Tier	Genomic Alteration	Relevant Therapies (In this cancer type)	Relevant Therapies (In other cancer type)	Clinical Trials
IA	ERBB2 exon 20 insertion	ado-trastuzumab emtansine	None	26
	erb-b2 receptor tyrosine kinase 2 Allele Frequency: 61.42%			

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 Varia	ants						
Gene	Amino Acid Change	Coding	Variant ID	Locus	Allele Frequency	Transcript	Variant Effect	Coverage
ERBB2	p.(E770_A771insAYV M)	c.2324_2325insATAC GTGATGGC	COSM20959	chr17:37880981	61.42%	NM_004448.3	nonframeshift Insertion	1965
JAK1	p.(=)	c.2199A>G		chr1:65310489	45.40%	NM_002227.3	synonymous	1987
ALK	p.(D1529E)	c.4587C>G		chr2:29416366	53.18%	NM_004304.4	missense	1854
ALK	p.(I1461V)	c.4381A>G		chr2:29416572	99.95%	NM_004304.4	missense	2000
ALK	p.(=)	c.3375C>A		chr2:29445458	50.68%	NM_004304.4	synonymous	1977
FGFR3	p.(=)	c.1206C>A		chr4:1806187	44.48%	NM_000142.4	synonymous	634
FGFR3	p.(=)	c.1953G>A		chr4:1807894	99.85%	NM_000142.4	synonymous	1996
PDGFRA	p.(=)	c.1701A>G		chr4:55141055	99.60%	NM_006206.5	synonymous	1995
FGFR4	p.(P136L)	c.407C>T		chr5:176517797	99.15%	NM_213647.2	missense	2000
RET	p.(=)	c.2307G>T		chr10:43613843	43.99%	NM_020975.4	synonymous	1996
RET	p.(=)	c.2712C>G		chr10:43615633	23.51%	NM_020975.4	synonymous	1999

Biomarker Descriptions

ERBB2 (erb-b2 receptor tyrosine kinase 2)

Background: The ERBB2 gene encodes the erb-b2 receptor tyrosine kinase 2, a member of the human epidermal growth factor receptor (HER) family. Along with ERBB2/HER2, EGFR/ERBB1/HER1, ERBB3/HER3, and ERBB4/HER4 make up the HER protein family¹. All ERBB/HER proteins encode transmembrane receptor tyrosine kinases. However, ERBB2/HER2 is an orphan receptor with no known ligand. ERBB2 preferentially binds other ligand bound ERBB/HER family members to form hetero-dimers resulting in the activation of ERBB2 tyrosine kinase activity and subsequent activation of the PI3K/AKT/MTOR and RAS/RAF/MAPK/ERK signaling pathways which promote cell proliferation, differentiation, and survival². Recurrent focal amplification of the ERBB2 gene leads to increased expression in several cancer types. ERBB2 overexpression in immortalized cell lines is oncogenic and leads to ERBB2 homo-dimerization and activation without ligand binding³.4.5.

Alterations and prevalence: ERBB2 gene amplification occurs in 10-20% of breast, esophageal, and gastric cancers, 5-10% of bladder, cervical, pancreas, and uterine cancers, and 1-5% of colorectal, lung, and ovarian cancers^{6,7,8,9,10,11,12,13}. Recurrent somatic activating mutations in ERBB2/HER2 occur at low frequencies (<1%) in diverse cancer types^{13,14,15}. In breast, bladder, and colorectal cancers,



Tel: 02-2875-7449

Date: 15 Oct 2020 3 of 8

Biomarker Descriptions (continued)

the most common recurrent ERBB2 activating mutations include kinase domain mutations L755S and V777L and the extracellular domain mutation S310F. In lung cancer, the most common recurrent ERBB2 activating mutations include in-frame exon 20 insertions, particularly Y772_A775dup.

Potential relevance: The discovery of ERBB2/HER2 as an important driver of breast cancer in 1987 led to the development of trastuzumab, a humanized monoclonal antibody with specificity to the extracellular domain of HER216,17. Trastuzumab18 was FDA approved for the treatment of HER2 positive breast cancer in 1998, and subsequently in HER2 positive metastatic gastric and gastroesophageal junction adenocarcinoma in 2010. Additional monoclonal antibody therapies have been approved by the FDA for HER2-positive breast cancer including pertuzumab19 (2012), a humanized monoclonal antibody that inhibits HER2 dimerization, and ado-trastuzumab emtansine²⁰ (2013), a conjugate of trastuzumab and a potent antimicrotubule agent. The combination of pertuzumab, trastuzumab, and a taxane is the preferred front-line regimen for HER2-positive metastatic breast cancer²¹. In addition to monoclonal antibodies, the small molecule inhibitor lapatinib²², with specificity for both EGFR and ERBB2, was FDA approved (2007) for the treatment of patients with advanced HER2-positive breast cancer who have received prior therapy including trastuzumab. In 2017, the FDA approved the use of neratinib23, an irreversible kinase inhibitor of EGFR, ERBB2/HER2, and ERBB4, for the extended adjuvant treatment of adult patients with early stage HER2-positive breast cancer. In 2020, the FDA approved neratinib²³ in combination with capecitabine for HER2-positive advanced or metastatic patients after two or more prior HER2-directed therapies. The vaccine, nelipepimut-S²⁴, was granted fast-track designation by the FDA (2016) in patients with low to intermediate HER2 expressing (IHC score 1+ or 2+) breast cancer. Additionally, fast-track designation was granted (2018) to the monoclonal antibody margetuximab²⁵ in patients with ERBB2 positive breast cancer previously treated with an anti-HER2 therapy as well as the novel bispecific antibody ZW25²⁶ (2019) in combination with standard chemotherapy for patients with HER2-overexpressing gastroesophageal adenocarcinoma (GEA). Certain activating mutations have been observed to impart sensitivity to neratinib, afatinib, lapatinib, and trastuzumab, or dacomitinib in early and ongoing clinical studies^{27,28,29,30,31}. Additionally, acquired HER2 mutations in estrogen receptor-positive (ER+) breast cancer have been shown to confer resistance to hormone therapy³². However, this was shown to be overcome by neratinib in combination with therapies targeting ER³².

Relevant Therapy Summary

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
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ERBB2 exon 20 insertion					
Relevant Therapy	FDA	NCCN	EMA	ESMO	Clinical Trials*
ado-trastuzumab emtansine	×		×	×	(II)
afatinib	×	×	×	×	(II)
anti-PD-L1 antibody, pyrotinib	×	×	×	×	(II)
neratinib	×	×	×	×	(II)
pertuzumab + trastuzumab	×	×	×	×	(II)
pertuzumab, trastuzumab	×	×	×	×	(II)
poziotinib	×	×	×	×	(II)
pyrotinib	×	×	×	×	(II)
sintilimab	×	×	×	×	(II)

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



Tel: 02-2875-7449

Date: 15 Oct 2020 4 of 8

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

ERBB2 exon 20 insertion (continued)

Relevant Therapy	FDA	NCCN	EMA	ESMO	Clinical Trials*
targeted therapy, chemotherapy	×	×	×	×	(II)
tarloxotinib	×	×	×	×	(II)
trastuzumab deruxtecan	×	×	×	×	(II)
trastuzumab, pertuzumab, chemotherapy	×	×	×	×	(II)
BDTX-189	×	×	×	×	(1/11)
CBT-502, anlotinib hydrochloride	×	×	×	×	(1/11)
DZD-9008	×	×	×	×	(1/11)
zotatifin	×	×	×	×	(1/11)
disitamab vedotin	×	×	×	×	(I)
neratinib, palbociclib, everolimus, trametinib	×	×	×	×	(l)
pirotinib	×	×	×	×	(l)
trastuzumab deruxtecan, pembrolizumab	×	×	×	×	(I)

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



Tel: 02-2875-7449

Date: 15 Oct 2020 5 of 8

Relevant Therapy Details

Current NCCN Information

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.

ERBB2 exon 20 insertion

ado-trastuzumab emtansine

Cancer type: Non-Small Cell Lung Cancer Variant class: ERBB2 mutation

NCCN Recommendation category: 2A

Population segment (Line of therapy):

Non-Small Cell Lung Cancer; Emerging biomarker in metastatic disease (Not specified)

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

afatinib

Variant class: ERBB2 mutation Cancer type: Non-Small Cell Lung Cancer

Summary:

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

"The NCCN NSCLC Panel does not recommend single-agent therapy with trastuzumab or afatinib (both for ERBB2 mutations), because response rates are lower and treatment is less effective when these agents are used for patients with ERBB2 mutations."

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

trastuzumab

Cancer type: Non-Small Cell Lung Cancer Variant class: ERBB2 mutation

Summary:

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

"The NCCN NSCLC Panel does not recommend single-agent therapy with trastuzumab or afatinib (both for ERBB2 mutations), because response rates are lower and treatment is less effective when these agents are used for patients with ERBB2 mutations."

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

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Tel: 02-2875-7449

Date: 15 Oct 2020 6 of 8

	Date. 13 Oct 2020	0 01 0
Signatures		
Testing Personnel:		
Laboratory Supervisor:		
Pathologist:		

Tel: 02-2875-7449

Date: 15 Oct 2020 7 of 8

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Date: 15 Oct 2020 8 of 8

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