

# REPORT SUMMARY

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Project ID: C21-M001-01002 Report No.: AA-21-04732\_ONC Date Reported: Nov 04, 2021

#### PATIENT AND SAMPLE INFORMATION

**PATIENT SPECIMEN ORDERING PHYSICIAN** 

Name: 林巧暄 Name: 洪君儀醫師 Type: FFPE tissue Gender: Female Date received: Oct 22, 2021 Facility: 臺北榮總 Date of Birth: Aug 28, 1992 Collection site: Bone Tel: 886-228712121 Patient ID: 46185813 Specimen ID: S11029141

Diagnosis: Osteosarcoma Lab ID: AA-21-04732

D/ID: NA

Address: 臺北市北投區石牌路二段 201 號

### **VARIANT(S) WITH CLINICAL RELEVANCE**

Only variant(s) with clinical significance are listed. See the "DETAILED TEST RESULTS" section for full details.

#### SINGLE NUCLEOTIDE AND SMALL INDEL VARIANTS

Not detected.

#### **COPY NUMBER VARIANTS (CNVS)**

Loss of heterozygosity (LOH) information was used to infer tumor cellularity. Copy number alteration in the tumor was determined based on 67% tumor purity.

Amplification (Conv. number > 8)

Amplification (copy number 2 8)						
Chr	Copy Number					
ND	ND	ND				

Homozygous deletion (Copy number=0)

ND

780 00 000 000 000	,				
Chr	Gene				
chr17	NF1				
Heterozygous deletion (Copy number=1)					
Chr Gene					

ND

ND, Not Detected

#### **TUMOR MUTATIONAL BURDEN (TMB) MICROSATELLITE INSTABILITY (MSI)**

Microsatellite stable (MSS) 1.2 muts/Mb

Muts/Mb, mutations per megabase

#### Note:

TMB was calculated by using the sequenced regions of ACTOnco®+ to estimate the number of somatic nonsynonymous mutations per megabase of all protein-coding genes (whole exome). The threshold for high mutation load is set at ≥ 7.5 mutations per megabase. TMB, microsatellite status and gene copy number deletion cannot be determined if calculated tumor purity is < 30%.

Variant Analysis:

醫藥資訊研究員 楊杭哲 博士 Hang-Che Yang Ph.D.

Sign Off

解剖病理專科醫師王業翰 Yeh-Han Wang M.D. 病解字第 000545 號

行動基因僅提供技術檢測服務及檢測報告,檢測結果之臨床解釋及相關醫療處置,請諮詢專業醫師。報告結果僅對此試驗件有效。

行動基因臨床分子醫學實驗室 台北市內湖區新湖二路 345 號 3F

Email: <a href="mailto:service@actgenomics.com">service@actgenomics.com</a> T: +886-2-2795-3660 | F: +886-2-2795-5016

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### THERAPEUTIC IMPLICATIONS

TARGETED THERAPIES			
Genomic Alterations	Therapies	Effect	
Level 3B			
NF1 Homozygous deletion Selumetinib sensi			
Level 4			
<b>NF1</b> Homozygous deletion	Everolimus, Trametinib	sensitive	
<b>NF1</b> Homozygous deletion	Afatinib, Erlotinib, Gefitinib, Cetuximab, Trastuzumab, Lapatinib, Vemurafenib	resistant	

 $<sup>^{\</sup>dagger}$  Refer to "ONGOING CLINICAL TRIALS" section for detailed trial information.

Note: Therapies associated with benefit or lack of benefit are based on biomarkers detected in this tumor and published evidence.

		——————————————————————————————————————		
Lev	/el	Description		
1	1	FDA-recognized biomarker predictive of response to an FDA approved drug in this indication		
2	2	Standard care biomarker (recommended as standard care by the NCCN or other expert panels) predictive of response to an FDA approved drug in this indication		
3	Α	Biomarkers that predict response or resistance to therapies approved by the FDA or professional societies for a different type of tumor		
	B Biomarkers that serve as inclusion criteria for clinical trials			
4	1	Biomarkers that show plausible therapeutic significance based on small studies, few case reports or preclinical studies		









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### **IMMUNE CHECKPOINT INHIBITORS (ICI) THERAPIES**

Genomic markers and alterations that are associated with response to ICI therapies

Positive Biomarker	Negative Biomarker
TMB-H: ND	EGFR aberration: ND
MSI-H: ND	MDM2/MDM4 amplification: ND
MMR biallelic inactivation: ND	STK11 biallelic inactivation: ND
PBRM1 biallelic inactivation: ND	PTEN biallelic inactivation: ND
SERPINB3/SERPINB4 mutation: ND	B2M biallelic inactivation: ND
<b>L</b>	JAK1/2 biallelic inactivation: ND

MMR, mismatch repair; ND, not detected

Note: Tumor non-genomic factors, such as patient germline genetics, PDL1 expression, tumor microenvironment, epigenetic alterations or other factors not provided by this test may affect ICI response.

#### **CHEMOTHERAPIES**

No genomic alterations detected in this tumor predicted to confer sensitivity or lack of benefit to chemotherapies.

HORMONAL THERAPIES					
Therapies	Genomic Alterations	Effect	Gene / Variant Level Evidence	Cancer Type	
Tamoxifen	<b>NF1</b> Homozygous deletion	less sensitive	Clinical	Breast cancer	

#### **OTHERS**

No genomic alterations detected in this tumor predicted to confer sensitivity or lack of benefit to other therapies.

#### Note:

Therapeutic implications provided in the test are based solely on the panel of 440 genes sequenced. Therefore, alterations in genes not covered in this panel, epigenetic and post-transcriptional and post-translational factors may also determine a patient's response to therapies. In addition, several other patient-associated clinical factors, including but not limited to, prior lines of therapies received, dosage and combinations with other therapeutic agents, patient's cancer types, sub-types, and/or stages, may also determine the patient's clinical response to therapies.









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### **VARIANT INTERPRETATION**

#### **NF1** Homozygous deletion

#### **Biological Impact**

The neurofibromin 1 (NF1) gene encodes a GTPase activating protein (GAP) which is an important negative regulator of the Ras cellular proliferation pathways<sup>[1][2][3][4]</sup>. Besides, NF1 also physically interacts with the N-terminal domain of focal adhesion kinase (FAK) and involves in the regulation of cell adhesion, growth, and other pathways<sup>[5][6]</sup>. NF1 is considered a classical haploinsufficient tumor suppressor gene with loss of one allele through inherited or acquired mutation may lead to reduced protein expression and is insufficient to execute normal cellular functions contributing to tumor development<sup>[7][8][9][10][11]</sup>. NF1 syndrome is a germline condition resulting in a predisposition to several types of cancer such as neurofibromas, melanoma, lung cancer, ovarian cancer, breast cancer, colorectal cancer, hematological malignancies<sup>[12][13][14]</sup>. Meanwhile, sporadic NF1 mutations have been observed in multiple cancer types<sup>[15]</sup>, including myelodysplastic syndromes, melanomas, colon cancer<sup>[16]</sup>, glioblastomas<sup>[17]</sup>, lung cancer<sup>[18]</sup>, ovarian cancer, and breast cancer<sup>[12]</sup>.

#### Therapeutic and prognostic relevance

In April 2020, the U.S. FDA has approved selumetinib for pediatric patients 2 years of age and older with neurofibromatosis type 1 (NF1) who have symptomatic, inoperable plexiform neurofibromas (PN).

A phase II trial (NCT02664935, NLMT) demonstrated that selumetinib in combination with docetaxel resulted in a confirmed ORR of 28.5% (4/14), durable clinical benefit (DCB) rate of 50% (7/14), and mPFS of 5.3 months in lung adenocarcinoma patients harboring NF1 loss<sup>[19]</sup>.

NF1 loss has been determined as an inclusion criterion for the trials evaluating selumetinib efficacies in lung cancer (NCT02664935) and NF1-associated tumors (NCT03326388).

A case study had reported that MEK inhibitor, trametinib, was effective in a treatment-refractory neurofibromatosis type I-associated glioblastoma<sup>[20]</sup>. Various preclinical data had also supported the activity of MEK<sup>[21][22][23][24]</sup> and mTOR<sup>[25][26]</sup> inhibitors in NF1-deficient tumors.

In an NGS-based study, patients harboring mutations in the mTOR pathway, including mTOR, TSC1, TSC2, NF1, PIK3CA, and PIK3CG responded to everolimus<sup>[27]</sup>.

NF1 depletion has been associated with drug resistance to RAF and EGFR inhibitors, tamoxifen, and retinoic acid<sup>[15][28]</sup>. For example, loss of NF1 was identified in patients with lung adenocarcinomas or colorectal cancer who presented resistance to anti-EGFR treatment, including erlotinib, gefitinib, afatinib, and cetuximab, respectively<sup>[29][30][31]</sup>. Notably, preclinical studies further revealed that the addition of a MEK inhibitor could restore the sensitivity to erlotinib<sup>[29]</sup>.





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In a liquid biopsy-based ctDNA profiling study of HER2-positive metastatic gastric cancer, NF1 loss (either induced by mutation or deletion) was suggested as a novel mechanism contributes to trastuzumab resistance. The cellbased study also showed that the trastuzumab resistance could be overcome with a combination of HER2 and MEK/ERK inhibitors<sup>[32]</sup>.

Loss of NF1 in patients with BRAF-mutated melanomas was suggested conferring resistance to BRAF inhibitors<sup>[33][34][35][36]</sup>







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### US FDA-APPROVED DRUG(S)

### **Everolimus (AFINITOR)**

Everolimus, a derivative of sirolimus, works as an inhibitor of mammalian target of rapamycin complex 1 (mTORC1) and blocks mTORC1-mediated downstream signals for cell growth, proliferation, and survival. Everolimus is developed and marketed by Novartis under the trade name AFINITOR.

### FDA Approval Summary of Everolimus (AFINITOR)

FDA Approvai Sullillar	of Everoninus (Arini Tok)
	Lung or gastrointestinal neuroendocrine tumor (Approved on 2016/02/26)
RADIANT-4 <sup>[37]</sup>	/-
NCT01524783	Everolimus vs. Placebo
	[PFS(M): 11 vs. 3.9]
	Breast cancer (Approved on 2012/07/20)
BOLERO-2 <sup>[38]</sup>	ER+/HER2-
NCT00863655	Everolimus + exemestane vs. Placebo + exemestane
	[PFS(M): 7.8 vs. 3.2]
	Pancreatic neuroendocrine tumor (Approved on 2011/05/05)
RADIANT-3 <sup>[39]</sup>	-
NCT00510068	Everolimus vs. Placebo
	[PFS(M): 11 vs. 4.6]
	Subependymal giant cell astrocytoma (Approved on 2010/10/29)
EXIST-1 <sup>[40]</sup>	-
NCT00789828	Everolimus vs. Placebo
	[ORR(%): 35.0]
	Renal cell carcinoma (Approved on 2009/05/30)
RECORD-1 <sup>[41]</sup>	-
NCT00410124	Everolimus vs. Placebo
	[PFS(M): 4.9 vs. 1.9]

#### **Selumetinib (KOSELUGO)**

Selumetinib is a kinase inhibitor. Selumetinib is developed and marketed by AstraZeneca under the trade name KOSELUGO.

#### FDA Approval Summary of Selumetinib (KOSELUGO)

	Plexiform neurofibromas (Approved on 2020/04/10)	
SPRINT	Neurofibromatosis type 1	
NCT01362803	Selumetinib	
	[ORR(%): 66.0]	

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### **Trametinib (MEKINIST)**

Trametinib is an anti-cancer inhibitor which targets MEK1 and MEK2. Trametinib is developed and marketed by GlaxoSmithKline (GSK) under the trade name MEKINIST.

#### FDA Approval Summary of Trametinib (MEKINIST)

TDA Approvar Summary of Transectinia (WEKINIST)				
	Anaplastic thyroid cancer (Approved on 2018/05/04)			
BRF117019 <sup>[42]</sup>	BRAF V600E			
NCT02034110	Dabrafenib + trametinib			
	[ORR(%): 61.0]			
	Non-small cell lung cancer (Approved on 2017/06/22)			
BRF113928 <sup>[43]</sup>	BRAF V600E			
NCT01336634	Trametinib + dabrafenib vs. Dabrafenib			
	[ORR(%): 63.0 vs. 27.0, DOR(M): 12.6 vs. 9.9]			
	Melanoma (Approved on 2014/01/10)			
COMBI-d <sup>[44]</sup>	BRAF V600E/K			
NCT01584648	Trametinib + dabrafenib vs. Dabrafenib + placebo			
	[PFS(M): 9.3 vs. 8.8]			
	Melanoma (Approved on 2013/05/29)			
METRIC <sup>[45]</sup>	BRAF V600E/K			
NCT01245062	Trametinib vs. Dacarbazine or paclitaxel			
	[PFS(M): 4.8 vs. 1.5]			

d=day; w=week; m=month

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行動基因臨床分子醫學實驗室 台北市內湖區新湖二路 345 號 3F

Email: <a href="mailto:service@actgenomics.com">service@actgenomics.com</a> T: +886-2-2795-3660 | F: +886-2-2795-5016

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### **ONGOING CLINICAL TRIALS**

Clinical trials shown below were selected by applying filters: study status, patient's diagnosis, intervention, location and/or biomarker(s). Please visit <a href="https://clinicaltrials.gov">https://clinicaltrials.gov</a> to search and view for a complete list of open available and updated matched trials.

No trial has been found.

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### **DETAILED TEST RESULTS**

### SINGLE NUCLEOTIDE AND SMALL INDEL VARIANTS

Gene	Chr	Exon	Accession Number	cDNA Change	Amino Acid Change	Coverage	Allele Frequency	COSMIC ID
AMER1	X	2	NM_152424	c.2047T>A	S683T	852	26.5%	-
ATRX	Х		NM_000489	c.189+7A>G	Splice region	1749	32.8%	-
BRCA1	17	7	NM_007294	c.509G>A	R170Q	740	81.9%	-
CDK6	7	2	NM_001259	c.224A>G	N75S	452	26.5%	-
CREBBP	16	6	NM_004380	c.1471C>T	P491S	516	58.3%	-
FAT1	4	19	NM_005245	c.10846G>A	V3616I	882	48.3%	COSM5648338
GNAQ	9	3	NM_002072	c.370T>A	F124I	1011	20.5%	-
GNAQ	9	3	NM_002072	c.380C>A	P127Q	1014	20.3%	COSM8792344
IRS2	13	1	NM_003749	c.1826C>T	A609V	960	97.4%	-
NOTCH2	1	13	NM_024408	c.2042T>A	1681N	820	78.3%	-
NOTCH4	6	3	NM_004557	c.316C>A	L106I	943	61.5%	-
RAD54L	1	16	NM_003579	c.1748T>C	I583T	821	62.1%	-

Mutations with clinical relevance are highlighted in red.





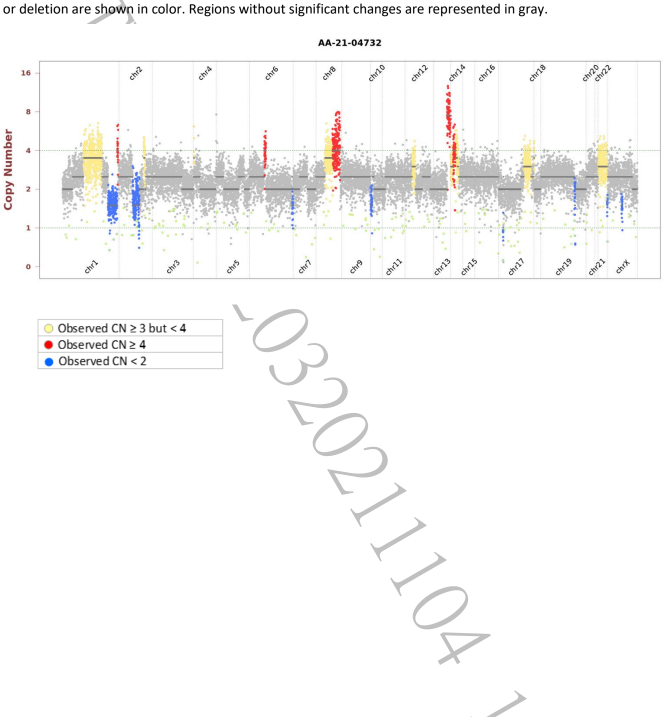




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### **COPY NUMBER VARIANTS (CNVS)**

Observed copy number (CN) for each evaluated position is shown on the y-axis. Regions referred to as amplification







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#### **HOTSPOT GENOTYPES**

Listed variants are biomarkers or hotspots that are recommended as standard care by the NCCN or other expert panels and not necessarily FDA-recognized for a particular indication. The genotypes have been manually checked to ensure sufficient coverage for each hotspot of the target gene.

Gene	Variant	<b>Genotype Detected</b>
BRAF	V600X	Not detected
EGFR	A763_Y764insFQEA, E709K, E709_T710delinsD, Exon 19 deletion, Exon 19 insertion, Exon 20 insertion, G719A/C/D/S, L747P, L833V, L858R, L861Q/R, S768I, T790M	Not detected
IDH2	R140Q, R172G/K/M/S	Not detected
KIT	A502_Y503dup, D419del, D579del, D816F/V/Y, D820A/E/G/Y, E554_I571del, E554_K558del, E554_V559del, Exon 11 mutation, F522C, H697Y, I563_L576del, I653T, K550_W557del, K558N, K558_E562del, K558_V559del, K558delinsNP, K642E, M552_W557del, N505I, N564_Y578del, N822H/I/K/Y, P551_M552del, P573_D579del, P577_D579del, P577_W582delinsPYD, P838L, Q556_K558del, T417_D419delinsI, T417_D419delinsRG, T574_Q575insTQLPYD, V530I, V555_L576del, V555_V559del, V559A/C/D/G, V559_V560del, V559del, V560D/G, V560del, V569_L576del, V654A, W557G/R, W557_K558del, Y553N, Y553_K558del, Y570H, Y578C	Not detected
KRAS	A146T/V/P, G12X, G13X, Q61X	Not detected
MET	D1028H/N/Y	Not detected
NRAS	G12X, G13X, Q61X	Not detected
PDGFRA	A633T, C450_K451insMIEWMI, C456_N468del, C456_R481del, D568N, D842I/V, D842_H845del, D842_M844del, D846Y, E311_K312del, G853D, H650Q, H845Y, H845_N848delinsP, I843del, N659K/R/S, N848K, P577S, Q579R, R560_V561insER, R748G, R841K, S566_E571delinsR, S584L, V469A, V536E, V544_L545insAVLVLLVIVIISLI, V561A/D, V561_I562insER, V658A, W559_R560del, Y375_K455del, Y555C, Y849C/S	Not detected
PIK3CA	C420R, E542K/V, E545A/D/G/K, H1047X, Q546E/R	Not detected

V600X= any mutation in the valine (V) at amino acid 600 being replaced by a different amino acid. G12X = any mutation in the glycine (G) at amino acid 12 being replaced by a different amino acid. G13X= any mutation in the glycine (G) at amino acid 13 being replaced by a different amino acid. Q61X = any mutation in the glutamine (Q) at amino acid 61 being replaced by a different amino acid. H1047X = any mutation in the histidine (H) at amino acid 1047 being replaced by a different amino acid.

Gene	Copy Number Detected
CDK4	2
EGFR	2
ERBB2	2
MET	3

Copy number ≥ 8 is considered amplification

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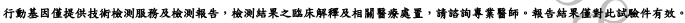
## **ACTOnco®** + Report

### Other known alterations that are associated with sensitivity, resistance, and toxicity to therapies.

Gene	Variant	<b>Genotype Detected</b>
AKT1	E17K	Not detected
ALK	C1156Y, D1203N, G1202R, L1152R, S1206Y, T1151_L1152insT	Not detected
BRAF	K601E, L597V/Q/R/S	Not detected
DPYD	D949V, I560S, splice-site mutation	Not detected
EGFR	A750P, C797S/Y, S492R	Not detected
ERBB2	V659E	Not detected
ESR1	D538G, E380Q, L469V, L536H/P/Q/R, S432L, S463P, V422del, V534E, Y537C/N/S	Not detected
FGFR3	G370C, G380R, K650E/N/R/M/T/Q, R248C, S249C, S371C, Y373C	Not detected
IDH1	R132C/G/H/L/Q/S	Not detected
MAP2K1	D67N, E203K, F53L, K57E/N, P124S, Q56P, Q56_V60del, R47Q, R49L, S222D	Not detected
PTEN	R130*/fs/G/L/P/Q	Not detected
TPMT	A154T, Y240C	Not detected

Gene	Copy Number Detected							
FGFR1	3							
MDM2	3							
MDM4	2							

Copy number ≥ 8 is considered amplification









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#### **TEST DETAILS**

#### **ABOUT ACTOnco®+**

The test is a next-generation sequencing (NGS)-based assay developed for efficient and comprehensive genomic profiling of cancers. This test interrogates coding regions of 440 genes associated with cancer treatment, prognosis and diagnosis. Genetic mutations detected by this test include small-scale mutations like single nucleotide variants (SNVs), small insertions and deletions (INDELs) (≤ 15 nucleotides) and large-scale genomic alterations like copy number variations (CNVs).

See ACTOnco®+ Gene List' Section for details of gene sequenced.

#### **DATABASE USED**

- Reference genome: human genome sequence hg19
- COSMIC v.92
- Genome Aggregation database r2.1.1
- ClinVar (version 20210208)
- ACT Genomics in-house database

#### **NEXT-GENERATION SEQUENCING (NGS) METHODS**

Extracted genomic DNA was amplified using four pools of primer pairs targeting coding exons of analyzed genes. Amplicons were ligated with barcoded adaptors. Quality and quantity of amplified library were determined using the fragment analyzer (AATI) and Qubit (Invitrogen). Barcoded libraries were subsequently conjugated with sequencing beads by emulsion PCR and enriched using Ion Chef system (Thermo Fisher Scientific) according to the Ion PI Hi-Q Chef Kit protocol (Thermo Fisher Scientific). Sequencing was performed on the Ion Proton or Ion S5 sequencer (Thermo Fisher Scientific).

Raw reads generated by the sequencer were mapped to the hg19 reference genome using the Ion Torrent Suite (version 5.10). Coverage depth was calculated using Torrent Coverage Analysis plug-in. Single nucleotide variants (SNVs) and short insertions/deletions (INDELs) were identified using the Torrent Variant Caller plug-in (version 5.10). The coverage was down-sampled to 4000. VEP (Variant Effect Predictor) (version 100) was used to annotate every variant using databases from Clinvar (version 20210208), COSMIC v.92 and Genome Aggregation database r2.1.1. Variants with coverage  $\geq$  25, allele frequency  $\geq$  5% and actionable variants with allele frequency  $\geq$  2% were retained.

This test provides uniform coverage of the targeted regions, enabling target base coverage at  $100x \ge 85\%$  with a mean coverage  $\ge 500x$ .

Variants reported in Genome Aggregation database r2.1.1 with > 1% minor allele frequency (MAF) were

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considered as polymorphisms. ACT Genomics in-house database was used to determine technical errors. Clinically actionable and biologically significant variants were determined based on the published medical literature.

The copy number variations (CNVs) were predicted as described below:

Amplicons with read counts in the lowest 5th percentile of all detectable amplicons and amplicons with a coefficient of variation ≥ 0.3 were removed. The remaining amplicons were normalized to correct the pool design bias. ONCOCNV (an established method for calculating copy number aberrations in amplicon sequencing data by Boeva et al., 2014) was applied for the normalization of total amplicon number, amplicon GC content, amplicon length, and technology-related biases, followed by segmenting the sample with a gene-aware model. The method was used as well for establishing the baseline of copy number variations from samples in ACT Genomics in-house database.

Tumor mutational burden (TMB) was calculated by using the sequenced regions of ACTOnco $^{\circ}$ + to estimate the number of somatic nonsynonymous mutations per megabase of all protein-coding genes (whole exome). The TMB calculation predicted somatic variants and applied a machine learning model with a cancer hotspot correction. TMB may be reported as "TMB-High", "TMB-Low" or "Cannot Be Determined". TMB-High corresponds to  $\geq$  7.5 mutations per megabase (Muts/Mb); TMB-Low corresponds to < 7.5 Muts/Mb. TMB is reported as "Cannot Be Determined" if the tumor purity of the sample is < 30%.

Classification of microsatellite instability (MSI) status is determined by a machine learning prediction algorithm. The change of a number of repeats of different lengths from a pooled microsatellite stable (MSS) baseline in > 400 genomic loci are used as the features for the algorithm. The final output of the results is either microsatellite Stable (MSS) or microsatellite instability high (MSI-H).

#### **STANDARD OPERATING PROCEDURES (SOPS)**

Standard operating procedures (SOPs) are shown below:

- AG2-QP-15 Specimen Management Procedure
- AG3-QP16-03 SOP of Cancer Cell DNA and RNA Extraction
- AG3-QP16-07 SOP of Nucleic Acid Extraction with QIAsymphony SP
- AG3-QP16-08 SOP of FFPE Nucleic Acid Extraction
- AG3-QP16-10 SOP of HE Staining
- AG3-QP16-13 SOP of Library Construction and Preparation
- AG3-QP16-17 SOP of DNA Quantification with Qubit Fluorometer
- AG3-QP16-20 SOP of CE-Fragment Analysis
- AG3-QP16-22 SOP of Variant Calling
- AG3-QP16-24 SOP of Ion Torrent System Sequencing Reaction
- AG3-QP16-26 SOP of Ion Chef Preparation





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## **ACTOnco® + Report**

- AG3-QP16-35 SOP of Variant Annotation
- AG3-QP16-96 SOP of Manual Inspection for SNVIndel Variant
- AG3-QP16-95 SOP of Manual Inspection for Copy Number Variant
- AG3-QP40-08 (02) Standard protocol for variant interpretation, curation and classification
- AG3-QP16-41 SOP of The user manual for clinical report system (CRS)

#### **LIMITATIONS**

This test does not provide information of variant causality and does not detect variants in non-coding regions that could affect gene expression. This report does not report polymorphisms and we do not classify whether a mutation is germline or somatic. Variants identified by this assay were not subject to validation by Sanger or other technologies.

#### **NOTES**

We do not exclude the possibility that pathogenic variants may not be reported by one or more of the tools and the parameters used.

#### **PATHOLOGY EVALUATION**

H&E-stained section No.: <u>S11029141</u>

Collection site: <u>Bone</u>

Examined by: <u>Dr. Yeh-Han Wang</u>

• Estimated neoplastic nuclei (whole sample): The percentage of viable tumor cells in total cells in the whole slide (%): 70%

The percentage of viable tumor cells in total cells in the encircled areas in the whole slide (%): 70%

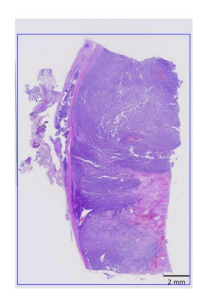
The percentage of necrotic cells (including necrotic tumor cells) in total cells in the whole slide (%): 0%

The percentage of necrotic cells (including necrotic tumor cells) in total cells in the encircled areas in the whole slide (%): 0%

Additional comment: NA

• Manual macrodissection: Not performed

The outline highlights the area of malignant neoplasm annotated by a pathologist.



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## **ACTOnco®+** Report

### **SPECIMEN PHOTO(S)**



• Collection date: <u>Sep 2021</u>

Facility retrieved: 臺北榮總

### **RUN QC**

Panel: <u>ACTOnco®+</u>
 Mean Depth: <u>659x</u>

Target Base Coverage at 100x: 92%







## 林巧暄

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## **ACTOnco®+ GENE LIST**

		1											
ABCB1*	AURKB	CBL	CDKN2B	E2F3	FAT1	GRIN2A	JAK2	MED12	<b>NOTCH4</b>	PMS1	RAD51D	SLCO1B3*	TNFRSF14
ABCC2*	AXIN1	CCNA1	CDKN2C	EGFR	FBXW7	GSK3B	JAK3	MEF2B	NPM1	PMS2	RAD52	SMAD2	TNFSF11
ABCG2*	AXIN2	CCNA2	CEBPA*	EP300	FCGR2B	GSTP1*	JUN*	MEN1	NQ01*	POLB	RAD54L	SMAD3	TOP1
ABL1	AXL	CCNB1	CHEK1	EPCAM	FGF1*	GSTT1*	KAT6A	MET	NRAS	POLD1	RAF1	SMAD4	TP53
ABL2	B2M	CCNB2	CHEK2	ЕРНА2	FGF10	HGF	KDM5A	MITF	NSD1	POLE	RARA	SMARCA4	ТРМТ*
ADAMTS1	BAP1	CCNB3	CIC	ЕРНА3	FGF14	HIF1A	KDM5C	MLH1	NTRK1	PPARG	RB1	SMARCB1	TSC1
ADAMTS13	BARD1	CCND1	CREBBP	ЕРНА5	FGF19*	HIST1H1C*	KDM6A	MPL	NTRK2	PPP2R1A	RBM10	SMO	TSC2
ADAMTS15	BCL10	CCND2	CRKL	ЕРНА7	FGF23	HIST1H1E*	KDR	MRE11	NTRK3	PRDM1	RECQL4	SOCS1*	TSHR
ADAMTS16	BCL2*	CCND3	CRLF2	ЕРНВ1	FGF3	HNF1A	KEAP1	MSH2	PAK3	PRKAR1A	REL	SOX2*	TYMS
ADAMTS18	BCL2L1	CCNE1	CSF1R	ERBB2	FGF4*	HR	КІТ	MSH6	PALB2	PRKCA	RET	SOX9	U2AF1
ADAMTS6	BCL2L2*	CCNE2	CTCF	ERBB3	FGF6	HRAS*	КМТ2А	MTHFR*	PARP1	PRKCB	RHOA	SPEN	UBE2A*
ADAMTS9	BCL6	ССПН	CTLA4	ERBB4	FGFR1	HSP90AA1	кмт2С	MTOR	PAX5	PRKCG	RICTOR	SPOP	UBE2K
ADAMTSL1	BCL9	CD19	CTNNA1	ERCC1	FGFR2	HSP90AB1	KMT2D	MUC16	PAX8	PRKCI	RNF43	SRC	UBR5
ADGRA2	BCOR	CD274	CTNNB1	ERCC2	FGFR3	HSPA4	KRAS	MUC4	PBRM1	PRKCQ	ROS1	STAG2	UGT1A1*
ADH1C*	BIRC2	CD58	CUL3	ERCC3	FGFR4	HSPA5	LCK	мис6	PDCD1	PRKDC	RPPH1	STAT3	USH2A
AKT1	BIRC3	CD70*	CYLD	ERCC4	FH	IDH1	LIG1	митүн	PDCD1LG2	PRKN	RPTOR	STK11	VDR*
AKT2	BLM	CD79A	CYP1A1*	ERCC5	FLCN	IDH2	LIG3	МҮС	PDGFRA	PSMB8	RUNX1	SUFU	VEGFA
АКТЗ	BMPR1A	CD79B	CYP2B6*	ERG	FLT1	IFNL3*	LMO1	MYCL	PDGFRB	PSMB9	RUNX1T1	SYK	VEGFB
ALDH1A1*	BRAF	CDC73	CYP2C19*	ESR1	FLT3	IGF1	LRP1B	MYCN	PDIA3	PSME1	RXRA	SYNE1	VHL
ALK	BRCA1	CDH1	CYP2C8*	ESR2	FLT4	IGF1R	LYN	MYD88	PGF	PSME2	SDHA	TAF1	WT1
AMER1	BRCA2	CDK1	CYP2D6	ETV1	FOXL2*	IGF2	MALT1	NAT2*	РНОХ2В*	PSME3	SDHB	TAP1	XIAP
APC	BRD4	CDK12	CYP2E1*	ETV4	FOXP1	IKBKB	MAP2K1	NBN	PIK3C2B	РТСН1	SDHC	TAP2	XPO1
AR	BRIP1	CDK2	CYP3A4*	EZH2	FRG1	IKBKE	MAP2K2	NEFH	PIK3C2G	PTEN	SDHD	ТАРВР	XRCC2
ARAF	BTG1*	CDK4	CYP3A5*	FAM46C	FUBP1	IKZF1	МАР2К4	NF1	РІКЗСЗ	PTGS2	SERPINB3	ТВХЗ	ZNF217
ARID1A	BTG2*	CDK5	DAXX	FANCA	GATA1	IL6	МАРЗК1	NF2	PIK3CA	PTPN11	SERPINB4	TEK	
ARID1B	ВТК	CDK6	DCUN1D1	FANCC	GATA2	IL7R	МАРЗК7	NFE2L2	РІКЗСВ	PTPRD	SETD2	TERT	
ARID2	BUB1B	CDK7	DDR2	FANCD2	GATA3	INPP4B	МАРК1	NFKB1	PIK3CD	PTPRT	SF3B1	TET1	
ASXL1	CALR	CDK8	DICER1	FANCE	GNA11	INSR	МАРК3	NFKBIA	PIK3CG	RAC1	SGK1	TET2	
ATM	CANX	CDK9	DNMT3A	FANCF	GNA13	IRF4	MAX	NKX2-1*	PIK3R1	RAD50	SH2D1A*	TGFBR2	
ATR	CARD11	CDKN1A	DOT1L	FANCG	GNAQ	IRS1	MCL1	NOTCH1	PIK3R2	RAD51	SLC19A1*	TMSB4X*	
ATRX	CASP8	CDKN1B	DPYD	FANCL	GNAS	IRS2*	MDM2	NOTCH2	PIK3R3	RAD51B	SLC22A2*	TNF	
AURKA	CBFB	CDKN2A	DTX1	FAS	GREM1	JAK1	MDM4	<b>NOTCH3</b>	PIM1	RAD51C	SLCO1B1*	TNFAIP3	

<sup>\*</sup>Analysis of copy number alteration not available.

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#### **DISCLAIMER**

#### **Legal Statement**

This test was developed by ACT Genomics and its performing characteristics were determined by ACT Genomics. This test result is to be used for clinical consultative purposes only and is not intended as a substitute for a clinical guidance of your doctor or another qualified medical practitioner. It should not be regarded as investigational or used for research.

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#### Treatment Decisions are the Responsibility of the Physician

Decisions on clinical care and treatment should be based on the independent medical judgment of the treating physician, taking into consideration all applicable information concerning the patient's condition, including physical examinations, information from other diagnostics tests and patient preferences, in accordance with the standard of care in a given community. A treating physician's decisions should not be based on a single test, such as this test, or the information contained in this report.

In terms of consulting a different treating physician, the patient must file an application and fulfill the listed criteria for ACT Genomics to provide the patient's report to the assigned physician. The report may not be copied or reproduced except in its totality.

#### Genetic Alterations and Drugs Not Presented in Ranked Order

In this report, neither any biomarker alteration nor any drug associated with a potential clinical benefit (or potential lack of clinical benefit), are ranked in order of potential or predicted efficacy.

#### **Level of Evidence Provided**

Drugs with a potential clinical benefit (or potential lack of clinical benefit) are evaluated for level of published evidence with at least one clinical efficacy case report or preclinical study. We endeavor to keep the information in the report up to date. However, customers must be aware that scientific understanding and technologies change over time, and we make no warranty as to the accuracy, suitability or currency of information provided in this report at any time.

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### 免責聲明

#### 法律聲明

本檢驗報告僅提供專業醫療參考,結果需經專業醫師解釋及判讀。基因突變資訊非必具備藥物或治療有效性指標,反之亦然。本檢驗報告提供之用藥指引不聲明或保證其臨床有效性,反之亦然。本基因檢測方法係由本公司研究開發,已經過有效性測試。

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#### 醫療決策需由醫師決定

任何治療與用藥需經由醫師在考慮病患所有健康狀況相關資訊包含健檢、其他檢測報告和病患意願後,依 照該地區醫療照護標準由醫師獨立判斷。醫師不應僅依據單一報告結果(例如本檢測或本報告書內容)做決策。

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藥物潛在臨床效益(或缺乏潛在臨床效益)的實證證據是依據至少一篇臨床療效個案報告或臨床前試驗做為評估。本公司盡力提供適時及準確之資料,但由於醫學科技之發展日新月異,本公司不就本報告提供的資料是否為準確、適宜或最新作保證。

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Email: <u>service@actgenomics.com</u> T: +886-2-2795-3660 | F: +886-2-2795-5016

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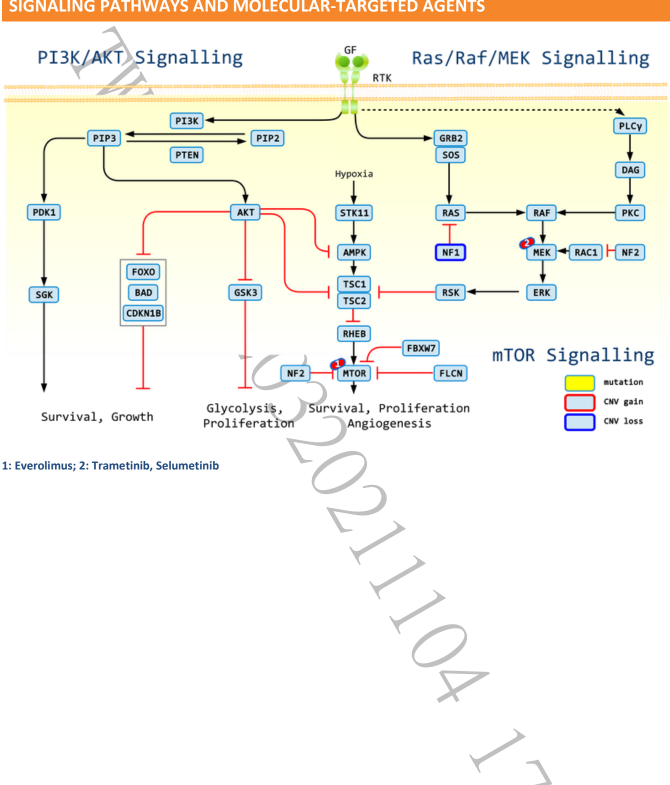




### 林巧暄

Project ID: C21-M001-01002 Report No.: AA-21-04732\_ONC Date Reported: Nov 04, 2021

### SIGNALING PATHWAYS AND MOLECULAR-TARGETED AGENTS







Project ID: C21-M001-01002 Report No.: AA-21-04732 ONC Date Reported: Nov 04, 2021

林巧暄

## ACTOnco® + Report

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行動基因僅提供技術檢測服務及檢測報告,檢測結果之臨床解釋及相關醫療處置,請諮詢專業醫師。報告結果僅對此試驗件有效。





Project ID: C21-M001-01002 Report No.: AA-21-04732\_ONC Date Reported: Nov 04, 2021

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## **ACTFusion™** Report

林巧暄

Project ID: C21-M001-01002 Report No.: AA-21-04732\_FUS Date Reported: Nov 04, 2021

### PATIENT SPECIMEN ORDERING PHYSICIAN

D/ID: NA

Name: 林巧暄 Gender: Female Date of Birth: Aug 28, 1992 Patient ID: 46185813 Diagnosis: Osteosarcoma Type: FFPE tissue
Date received: Oct 22, 2021
Collection site: Bone
Specimen ID: S11029141
Lab ID: AA-21-04732

Name: 洪君儀醫師 Facility: 臺北榮總 Tel: 886-228712121

Address: 臺北市北投區石牌路二段 201 號

#### **ABOUT ACTFusion**TM

The test is a next-generation sequencing (NGS) based in vitro diagnostic assay to detect fusion transcripts of 13 genes, including ALK, BRAF, EGFR, FGFR1, FGFR2, FGFR3, MET, NRG1, NTRK1, NTRK2, NTRK3, RET, and ROS1.

### **VARIANT(S) WITH CLINICAL RELEVANCE**

#### **FUSION RESULTS**

No fusion gene detected in this sample.

**Variant Analysis:** 

醫藥資訊研究員 楊杭哲 博士 Hang-Che Yang Ph.D. hay

Sign Off

解剖病理專科醫師王業翰 Yeh-Han Wang M.D. 病解字第 000545 號

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行動基因臨床分子醫學實驗室 台北市內湖區新湖二路 345 號 3F

Email: <u>service@actgenomics.com</u> | T: +886-2-2795-3660 | F: +886-2-2795-5016

AG4-QP4006-04(01)





## **ACTFusion**™ Report

林巧暄

Project ID: C21-M001-01002 Report No.: AA-21-04732\_FUS Date Reported: Nov 04, 2021

### THERAPEUTIC IMPLICATIONS

**TARGETED THERAPIES** 

Not Applicable.

### **VARIANT INTERPRETATION**

Not Applicable.

## **US FDA-APPROVED DRUG(S)**

Not Applicable.



行動基因臨床分子醫學實驗室 台北市內湖區新湖二路 345 號 3F Email: <u>service@actgenomics.com</u> | T: +886-2-2795-3660| F: +886-2-2795-5016

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# **ACTFusion**<sup>™</sup> Report

林巧暄

Project ID: C21-M001-01002 Report No.: AA-21-04732\_FUS Date Reported: Nov 04, 2021

### **ONGOING CLINICAL TRIAL(S)**

Clinical trials shown below were selected by applying filters: study status, patient's diagnosis, intervention, location and/or biomarker(s). Please visit <a href="https://clinicaltrials.gov">https://clinicaltrials.gov</a> to search and view for a complete list of open available and updated matched trials.

No trial has been found.

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Email: <u>service@actgenomics.com</u> | T: +886-2-2795-3660| F: +886-2-2795-5016

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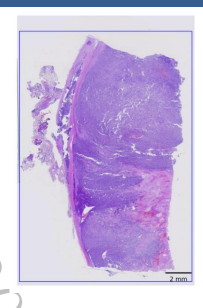
### **ACTFusion™ GENE LIST**

ALK	BRAF	EGFR	FGFR1	FGFR2	FGFR3	MET	NRG1
NTRK1	NTRK2	NTRK3	RET	ROS1	-	-	-

#### **TEST DETAILS**

### SPECIMEN RECEIVED





H&E-stained section No.: <u>S11029141</u>

• Collection date: Sep 2021

Collection site: <u>Bone</u>

Facility retrieved: 臺北榮總

Examined by: <u>Dr. Yeh-Han Wang</u>

• Estimated neoplastic nuclei (whole sample): The percentage of viable tumor cells in total cells in the whole slide (%): 70%

The percentage of viable tumor cells in total cells in the encircled areas in the whole slide (%): 70%

The percentage of necrotic cells (including necrotic tumor cells) in total cells in the whole slide (%): 0%

The percentage of necrotic cells (including necrotic tumor cells) in total cells in the encircled areas in the whole slide (%): 0%

Additional comment: NA

• Manual macrodissection: Not performed

The outline highlights the area of malignant neoplasm annotated by a pathologist.

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#### **NEXT-GENERATION SEQUENCING (NGS) METHODS**

The extracted RNA was reverse-transcribed and subjected to library construction. The quality and quantity of the amplified library was determined using the fragment analyzer (AATI) and Qubit (Invitrogen). Sequencing was performed on the Ion 540<sup>™</sup> Chip/ Ion 550<sup>™</sup> Chip / Ion P1<sup>™</sup> Chip and Ion GeneStudio<sup>™</sup> S5 Prime System / Ion Proton<sup>™</sup> System (Life Technologies). All assays were performed in accordance with ACT Genomics testing SOPs.

Data processing and statistical analysis for the identification of relevant fusions was performed using in-house fusion calling pipeline with default parameter setting. The four internal controls for the purpose of monitoring the overall sequencing quality of the sample were built into the assay, including CHMP2A, RABA7A, GPI, and VCP. Amplification of these genes using gene specific primers was performed, and the sequencing results were applied to the analysis pipeline to assess RNA quality. The inability of the software to detect these genes was considered a run failure. To ensure optimal sequencing quality for variant analysis, all samples had to meet the following sample quality control (QC) criteria: 1) Average unique RNA Start Sites (SS) per control Gene Specific Primer 2 (GSP 2)  $\geq$  10 (default), and 2) Total reads after sequencing  $\geq$  500,000 (recommended).

Samples passed the sample QC would be subjected to the fusion analysis pipeline for fusion transcript calling. Briefly, the analysis pipeline aligned sequenced reads to a reference genome, identified regions that map to noncontiguous regions of the genome, and applied filters to exclude probable false-positive events and annotate previously characterized fusion events. A minimum of 5 reads with 3 unique sequencing start sites that cross the breakpoints was set as the cutoff value to indicate strong evidence of fusions. RNA fusions would need to be in frame in order to generate productive transcripts. In addition, databases with details for documented fusions were used to authenticate the fusion sequence identified. Known fusions were queried using Quiver Gene Fusion Database, a curated database owned and maintained by ArcherDX. In summary, samples with detectable fusions had to meet the following criteria: 1) Number of unique start sites (SS) for the GSP2  $\geq$  3. 2) Number of supporting reads spanning the fusion junction  $\geq$  5. 3) Percentage of supporting reads spanning the fusion junction  $\geq$  10%. 4) Fusions annotated in Quiver Gene Fusion Database.

#### **DATABASE USED**

Quiver Gene Fusion Database version 5.1.18

#### **LIMITATIONS**

This test has been designed to detect fusions in 13 genes sequenced. Therefore, fusion in genes not covered by this test would not be reported. For novel fusions detected in this test, Sanger sequencing confirmation is recommended if residue specimen is available.





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#### **STANDARD OPERATING PROCEDURES (SOPS)**

Standard operating procedures (SOPs) are shown below:

- AG2-QP-15 Specimen Management Procedure
- AG3-QP16-08 SOP of FFPE Nucleic Acid Extraction
- AG3-QP16-10 SOP of HE Staining
- AG3-QP16-17 SOP of DNA Quantification with Qubit Fluorometer
- AG3-QP16-20 SOP of CE-Fragment Analysis
- AG3-QP16-24 SOP of Ion Torrent System Sequencing Reaction
- AG3-QP16-26 SOP of Ion Chef Preparation
- AG3-QP40-08 (02) Standard protocol for variant interpretation, curation and classification
- AG3-QP16-94 (01) SOP of ACTFusion v3 Library Construction and Preparation
- AG3-QP16-36(02) SOP of Fusion Gene Detection
- AG3-QP16-41 SOP of The user manual for clinical report system (CRS)

#### **RUN QC**

- Panel: <u>ACTFusion™</u>
- Total reads: 772638
- Average unique RNA Start Sites per control GSP2: <u>156</u>

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#### **DISCLAIMER**

#### **Legal Statement**

This test was developed by ACT Genomics and its performing characteristics were determined by ACT Genomics. This test result is to be used for clinical consultative purposes only and is not intended as a substitute for a clinical guidance of your doctor or another qualified medical practitioner. It should not be regarded as investigational or used for research.

The detection of genomic alterations does not necessarily indicate pharmacologic effectiveness (or lack thereof) of any drug or treatment regimen; the detection of no genomic alteration does not necessarily indicate lack of pharmacologic effectiveness (or effectiveness) of any drug or treatment regimen.

#### Treatment Decisions are the Responsibility of the Physician

Decisions on clinical care and treatment should be based on the independent medical judgment of the treating physician, taking into consideration all applicable information concerning the patient's condition, including physical examinations, information from other diagnostics tests and patient preferences, in accordance with the standard of care in a given community. A treating physician's decisions should not be based on a single test, such as this test, or the information contained in this report.

In terms of consulting a different treating physician, the patient must file an application and fulfill the listed criteria for ACT Genomics to provide the patient's report to the assigned physician. The report may not be copied or reproduced except in its totality.

#### **Genetic Alterations and Drugs Not Presented in Ranked Order**

In this report, neither any biomarker alteration nor any drug associated with a potential clinical benefit (or potential lack of clinical benefit), are ranked in order of potential or predicted efficacy.

#### **Level of Evidence Provided**

Drugs with a potential clinical benefit (or potential lack of clinical benefit) are evaluated for level of published evidence with at least one clinical efficacy case report or preclinical study. We endeavor to keep the information in the report up to date. However, customers must be aware that scientific understanding and technologies change over time, and we make no warranty as to the accuracy, suitability or currency of information provided in this report at any time.

#### No Guarantee of Clinical Benefit

This report makes no promises or guarantees about the effectiveness of a particular drug or any treatment procedure in any disease or in any patient. This report also makes no promises or guarantees that a drug without an association of reportable genomic alteration will, in fact, provide no clinical benefit.

#### Liability

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### 免責聲明

#### 法律聲明

本檢驗報告僅提供專業醫療參考,結果需經專業醫師解釋及判讀。基因突變資訊非必具備藥物或治療有效性指標,反之亦然。本檢驗報告提供之用藥指引不聲明或保證其臨床有效性,反之亦然。本基因檢測方法係由本公司研究開發,已經過有效性測試。

本檢驗報告非經本公司許可,不得私自變造、塗改,或以任何方式作為廣告及其他宣傳之用途。 本公司於提供檢驗報告後,即已完成本次契約義務,後續之報告解釋、判讀及用藥、治療,應自行尋求相關 專業醫師協助,若需將報告移件其他醫師,本人應取得該醫師同意並填寫移件申請書,主動告知行動基因, 行動基因僅能配合該醫師意願與時間提供醫師解說。

#### 醫療決策需由醫師決定

任何治療與用藥需經由醫師在考慮病患所有健康狀況相關資訊包含健檢、其他檢測報告和病患意願後,依 照該地區醫療照護標準由醫師獨立判斷。醫師不應僅依據單一報告結果(例如本檢測或本報告書內容)做決策。

#### 基因突變與用藥資訊並非依照有效性排序

本報告中列出之生物標記變異與藥物資訊並非依照潛在治療有效性排序。

#### 證據等級

藥物潛在臨床效益(或缺乏潛在臨床效益)的實證證據是依據至少一篇臨床療效個案報告或臨床前試驗做為評估。本公司盡力提供適時及準確之資料,但由於醫學科技之發展日新月異,本公司不就本報告提供的資料是否為準確、適宜或最新作保證。

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## **REFERENCES**

Not Applicable.

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