Project ID: C22-M001-01203 Report No.: AA-22-02113_ONC Date Reported: May 05, 2022

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PATIENT		
Name: 陳翠珠	Patient ID: 38194115	
Date of Birth: Jun 04, 1959	Gender: Female	
Diagnosis: Pancreatic cancer		
ORDERING PHYSICIAN		
Name: 周德盈醫師/姜乃榕醫師	Tel: 886-228712121	
Facility: 臺北榮總		
Address: 臺北市北投區石牌路二段 201 號		
SPECIMEN		
Specimen ID: S11170160A Collection site: Pancreas	Type: FFPE tissue	
Date received: Apr 22, 2022 Lab ID: AA-22-02113	D/ID: NA	

ABOUT ACTORCO®4

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 alterations (CNAs). The test also includes an RNA test, detecting fusion transcripts of 13 genes.

SUMMARY FOR ACTIONABLE VARIANTS VARIANTS/BIOMARKERS WITH EVIDENCE OF CLINICAL SIGNIFICANCE

Genomic	Probable Effects in F	Probable Sensitive in Other		
Alterations/Biomarkers	Sensitive Resistant		Cancer Types	
Not detected				

VARIANTS/BIOMARKERS WITH POTENTIAL CLINICAL SIGNIFICANCE

Genomic Alterations/Biomarkers	Possibly Sensitive	Possibly Resistant
BRAF V600_K601delinsE	Selumetinib	Vemurafenib
FGFR1 Amplification	Erdafitinib, Infigratinib, Lenvatinib, Pazopanib, Ponatinib, Regorafenib, Sunitinib	Palbociclib, Ribociclib

Note:

- The above summary tables present genomic variants and biomarkers based on the three-tiered approach proposed by US FDA for reporting tumor profiling NGS testing. "Variants/biomarkers with evidence of clinical significance" refers to mutations that are widely recognized as standard-of-care biomarkers (FDA level 2/AMP tier 1). "Variants/biomarkers with potential clinical significance" refers to mutations that are not included in the standard of care but are informational for clinicians, which are commonly biomarkers used as inclusion criterial for clinical trials (FDA level 3/AMP tier 2).
- The therapeutic agents and possible effects to a given drug are based on mapping the variants/biomarkers with ACT Genomics clinical knowledge database. The mapping results only provide information for reference, but not medical recommendation.
- Please refer to corresponding sections for more detailed information about genomic alteration and clinical relevance listed above.





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TESTING RESULTS

VARIANT(S) WITH CLINICAL RELEVANCE

- Single Nucleotide and Small InDel Variants

Gene	Amino Acid Change	Allele Frequency
BRAF	V600_K601delinsE	31.9%
TP53	Splice donor	42.9%

- Copy Number Alterations

Chromosome Gene		Variation	Copy Number
Chr18	SMAD4	Heterozygous deletion	1
Chr19	STK11	Heterozygous deletion	1
Chr3	BAP1	Heterozygous deletion	1
Chr8	FGFR1	Amplification	8

- Fusions

Fusion Gene & Exon	Transcript ID
	No fusion gene detected in this sample

- Immune Checkpoint Inhibitor (ICI) Related Biomarkers

Biomarker	Results	
Tumor Mutational Burden (TMB)	1.9 muts/Mb	
Microsatellite Instability (MSI)	Microsatellite stable (MSS)	

Note:

- Variant(s) enlisted in the SNV table may currently exhibit no relevance to treatment response prediction. Please refer to INTERPRETATION for more biological information and/or potential clinical impacts of the variants.
- Loss of heterozygosity (LOH) information was used to infer tumor cellularity. Copy number alteration in the tumor was determined based on 43% tumor purity.
- For more therapeutic agents which are possibly respond to heterozygous deletion of genes listed above, please refer to APPENDIX for more information.
- 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%.





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THERAPEUTIC IMPLICATIONS

TARGETED THERAPIES

Genomic Alterations	Therapies	Effect			
Level 3B					
BRAF V600_K601delinsE	Selumetinib	sensitive			
FGFR1 Amplification	Erdafitinib, Infigratinib, Ponatinib, Regorafenib, Sunitinib	sensitive			
Level 4					
FGFR1 Amplification	Lenvatinib, Pazopanib	sensitive			
BRAF V600_K601delinsE	Vemurafenib	resistant			
FGFR1 Amplification	Palbociclib, Ribociclib	resistant			

Therapies associated with benefit or lack of benefit are based on biomarkers detected in this tumor and published evidence in professional guidelines or peer-reviewed journals.

Level	Description
1	FDA-recognized biomarkers predictive of response or resistance to FDA approved drugs in this indication
2	Standard care biomarkers (recommended by the NCCN guideline) predictive of response or resistance to FDA approved drugs in this indication
ЗА	Biomarkers predictive of response or resistance to therapies approved by the FDA or NCCN guideline in a different cancer type
3B	Biomarkers that serve as inclusion criteria for clinical trials (minimal supportive data required)
4	Biomarkers that show plausible therapeutic significance based on small studies, few case reports, or preclinical studies





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IMMUNE CHECKPOINT INHIBITORS (ICIs)

No genomic alterations detected to confer sensitivity or lack of benefit to immune checkpoint therapies.

- Other Biomarkers with Potential Clinical Effects for ICIs

Genomic Alterations	Potential Clinical Effects
	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

Genomic Alterations	Therapies	Effect	Level of Evidence	Cancer Type
FGFR1 Amplification	Letrozole	Resistant	Clinical	Estrogen-receptor positive breast cancer
	Tamoxifen	Resistant	Preclinical	Breast cancer

OTHERS

Pharmacogenomic implication

Gene	Detection Site	Genotype	Drug Impact	Level of Evidence*
UGT1A1	rs4148323	AG	Irinotecan-based regimens	Level 1B

Clinical Interpretation:

Patients with the AG genotype and cancer who are treated with irinotecan-based regimens may have an increased risk of diarrhea and neutropenia as compared to patients with the GG genotype, or a decreased risk of diarrhea and neutropenia compared to patients with the AA genotype. Other genetic and clinical factors may also influence a patient's risk of diarrhea and neutropenia.

Level 1A: Clinical annotations describe variant-drug combinations that have variant-specific prescribing guidance available in a current clinical guideline annotation or an FDA-approved drug label annotation.

Level 1B: Clinical annotations describe variant-drug combinations with a high level of evidence supporting the association but no variant-specific prescribing guidance in an annotated clinical guideline or FDA drug label.

Level 2A: Variants in Level 2A clinical annotations are found in PharmGKB's Tier 1 Very Important Pharmacogenes (VIPs). These variants are in known pharmacogenes, implying causation of drug phenotype is more likely.

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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^{*} Level of evidence was defined by PharmGKB (https://www.pharmgkb.org/page/clinAnnLevels)

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VARIANT INTERPRETATION

BRAF V600 K601delinsE

Biological Impact

BRAF is a serine/threonine kinase that belongs to the RAF family. The protein plays an essential role in the regulation of mitogen-activated protein kinase (MAPK) cascade, which affects a range of cellular response including cell division, differentiation, and secretion[1][2]. Mutations in the BRAF gene, most commonly the V600 residue, are the most frequently identified oncogenic mutations in melanomas, and have been identified in several types of cancers including non-Hodgkin lymphoma, thyroid cancers, non-small cell lung carcinoma, hairy cell leukemia, glioma, gastrointestinal stromal tumor, and colorectal cancers (CRCs)[3][4]. Of note, in the vast majority of cases, BRAF mutations are nonoverlapping with other oncogenic mutations (e.g., NRAS mutations, KIT mutations, etc.) found in melanoma. V600E has been determined to be an activating mutation, which results in enhanced BRAF kinase activity and constitutive activation of downstream MEK/ERK signaling cascade^{[5][6]}.

BRAF V600 K601delinsE results in a deletion of 2 amino acids in the protein kinase domain of the BRAF protein from amino acids 600 to 601, combined with the insertion of a glutamate at the same site (UniProtKB). BRAF V600 K601delinsE has been identified in papillary thyroid cancer and NSCLC, and functionally characterized as an activating mutation that enhanced MEK/ERK signaling pathway and anchorage-dependent colony formation in vitro[7][8].

Therapeutic and prognostic relevance

A case report demonstrated that a NSCLC patient harboring BRAF V600_K601delinsE mutation showed disease progression after vemurafenib treatment[9].

The NCCN guidelines for central nervous system cancers recommended selumetinib for pilocytic astrocytoma patients with BRAF fusion or BRAF V600E activating mutation. BRAF activating mutations have been determined as an inclusion criterion for the trials evaluating selumetinib efficacies in cancers (NCT01089101, NCT00888134, NCT00866177, and NCT00936221).

TP53 Splice donor

Biological Impact

TP53 encodes the p53 protein, a crucial tumor suppressor that orchestrates essential cellular processes including cell cycle arrest, senescence and apoptosis[10]. TP53 is a proto-typical haploinsufficient gene, such that loss of a single copy of TP53 can result in tumor formation[11].

TP53 c.993+1G>A is a variant located at the splice donor region, which may result in the exon skipping.

Therapeutic and prognostic relevance

Despite having a high mutation rate in cancers, there are currently no approved targeted therapies for TP53 mutations. A phase II trial demonstrated that Wee1 inhibitor (AZD1775) in combination with carboplatin was well tolerated and showed promising anti-tumor activity in TP53-mutated ovarian cancer refractory or resistant (< 3 months) to standard first-line therapy (NCT01164995)[12].

In a retrospective study (n=19), advanced sarcoma patients with TP53 loss-of-function mutations displayed improved progression-free survival (208 days versus 136 days) relative to patients with wild-type TP53 when treated with pazopanib[13]. Results from another Phase I trial of advanced solid tumors (n=78) demonstrated that TP53 hotspot mutations are associated with better clinical response to the combination of pazopanib and vorinostat[14].

Advanced solid tumor and colorectal cancer patients harboring a TP53 mutation have been shown to be more sensitive to bevacizumab when compared with patients harboring wild-type TP53^{[15][16][17]}. In a pilot trial (n=21), TP53-negative





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breast cancer patients demonstrated increased survival following treatment with bevacizumab in combination with chemotherapy agents, Adriamycin (doxorubicin) and Taxotere (docetaxel)[18]. TP53 mutations were correlated with poor survival of advanced breast cancer patients receiving tamoxifen or primary chemotherapy[19][20]. In a retrospective study of non-small cell lung cancer (NSCLC), TP53 mutations were associated with high expression of VEGF-A, the primary target of bevacizumab, offering a mechanistic explanation for why patients exhibit improved outcomes after bevacizumab treatment when their tumors harbor mutant TP53 versus wild-type TP53[21].

BAP1 Heterozygous deletion

Biological Impact

Breast cancer type 1 susceptibility protein (BRCA1)-associated protein (BAP1) encodes an enzyme with ubiquitin carboxyl hydrolase activity involved in the regulation of cell cycle, transcription, and double-strand DNA repair[22][23][24]. BAP1 acts as a tumor suppressor by forming a complex with BRCA1^[25]. BAP1 is a haploinsufficient tumor suppressor with one copy loss may lead to weak protein expression and is related to renal cell carcinoma (RCC)[26]. Inactivating mutations of BAP1 were frequently observed in uveal melanoma with high metastatic risk, malignant mesothelioma other carcinoma types, including a subtype of renal cell carcinoma cholangiocarcinoma^{[23][27][28][29][30][31][32]}

Therapeutic and prognostic relevance

In a Phase II trial (MiST1; NCT03654833), rucaparib demonstrated manageable toxicity and clinical activity in patients with relapsed malignant mesothelioma that were negative for BAP1 (n=23), BRCA1 (n=13), or both (n=10), resulting in a 12-week disease control rate (DCR) of 58% (15/26), a 24-week DCR of 23% (6/26), and an objective response rate of 11.5% (3/26)[33]. The loss of BAP1 was shown to be associated with increased sensitivity to PARP inhibitor, olaparib, in renal cell carcinoma (RCC)[29] and mesothelioma cell lines[34]. However, no difference in sensitivity to the PARP inhibitor niraparib (MK4827) was observed between BAP1-mutant and wild-type mesothelioma cell[23]. BAP1 deficiency was also linked to a high tumor grade and was correlated with metastasis development in uveal melanoma[27].

An open-label, non-randomized, Phase II study (NCT03207347) has been initiated, aimed at investigating the use of niraparib in mesothelioma, uveal melanoma, renal cell carcinoma, and cholangiocarcinoma patients with tumors known to have mutations in BAP1 and other selected DNA double-strand break repair pathway genes. BAP1 loss of function mutation has been selected as an inclusion criteria for the trial examining olaparib in urothelial cancer (NCT03375307) and malignant mesothelioma (NCT04515836).

FGFR1 Amplification

Biological Impact

The fibroblast growth factor receptor 1 (FGFR1) gene encodes a receptor tyrosine kinase that plays crucial roles in cellular proliferation, survival, migration and angiogenesis[35][36]. Several studies have demonstrated that FGFR1 amplification correlates with FGFR1 overexpression[37][38][39][40][41][42]. Overexpression of FGFR1 has also been shown to enhance both ligand-dependent, and independent activation of downstream signaling pathways such as the phosphoinositide-3 kinase (PI3K) and the extracellular signal-regulated kinase 1/2 (ERK1/2) cascades[43][44][45]. Amplification of FGFR1 has been associated with early relapse, and poor survival, specifically in ER+ breast cancer^{[43][46]}, and may be associated with progression of breast cancer from in situ-to-invasive transition^[47].

FGFR1 amplifications have been reported in various types of cancer, including lung cancer^[48], breast cancer^[43], oral squamous cell carcinoma (OSCC)[49], prostate cancer[50], and esophageal cell carcinoma[51]. Besides, activating mutations (C381R and N330I) have been identified in giant cell lesions of the jaw^[52].

Therapeutic and prognostic relevance

Non-selective TKI-targeting inhibitors such as pazopanib, regorafenib, and ponatinib are multi-kinase inhibitors with





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inhibitory activities towards FGFR1[53][54].

To date, Erdafitinib (BALVERSATM), is the first and only pan-FGFR kinase inhibitor approved by U.S. FDA, for the treatment of patients with locally advanced or metastatic bladder cancer with FGFR3 mutations or FGFR2/FGFR3 fusions. Addition of the erdafitinib to palbociclib/fulvestrant induced complete responses of FGFR1-amplified/ER+ patient-derived-xenografts^[55].

A case report of a patient with HR+, HER2- breast cancer harboring FGFR1 amplification responded well to pazopanib^[56].

FGFR1 amplification has been selected as an inclusion criteria for the trial examining erdafitinib, ponatinib, regorafenib, sunitinib, and infigratinib efficacies in multiple tumor types (NCT03390504, NCT03473743, NCT03238196, NCT02272998, NCT02795156, NCT02693535, NCT04233567, NCT02150967).

Several small molecule FGFR inhibitors such as AZD-4547 and NVP-BGJ398 (Infigratinib) are under clinical evaluation, although mainly in the early stages of trials^[57]. Infigratinib has shown antitumor activity and manageable safety profile in patients with a variety of solid tumors, including FGFR1-amplified squamous cell lung cancer (sqNSCLC) and FGFR3-mutant bladder/urothelial cancers^[58]. Meanwhile, Dovitinib, a potent FGFR inhibitor, in combination with fulvestrant showed promising clinical activity in the FGF pathway-amplified postmenopausal patients with HR+, HER2-advanced breast cancer^[59].

In ER-positive breast cancer, FGFR1 amplification has been implicated as an acquired mechanism of resistance to endocrine therapies^[60], such as letrozole, 4-hydroxytamoxifen, and anastrozole-containing regimen^{[61][43][62]}. Besides, FGFR1/2 amplification or activating mutations were detected in ctDNA from post-progression ER-positive breast cancer patients after the fulvestrant plus palbociclib treatment. According to the subgroup analysis from MONALEESA-2 clinical trial, ER-positive breast cancer patients with FGFR1 amplification exhibited a shorter progression-free survival when treated with letrozole plus ribociclib^[55].

Meanwhile, in non-small cell lung carcinoma (NSCLC), FGFR1 is considered as an alternative acquired mechanism of resistance to EGFR tyrosine kinase inhibitors^[63]. For example, upregulated FGFR1-FGF2 autocrine loop was identified in a gefitinib-resistant cell model^[64], and focal FGFR1 amplification was observed in an NSCLC patient who developed resistance to osimertinib treatment^[65].

The BOLERO-2 clinical trial (everolimus plus exemestane) suggested that FGFR1 amplification and CCND1 amplification may be correlated with lessened progression-free survival (PFS) with the mTOR inhibitor everolimus [66][67].

In preclinical study, thyroid cancer cell with FGFR1 amplification is sensitive to lenvatinib treatment^{[68][69]}. Ponatinib, a multi-targeted tyrosine kinase inhibitor, demonstrated anti-proliferative activity in lung cancer, breast cancer, and Ewing's sarcoma cells overexpressing FGFR1^{[70][63][71]}.

SMAD4 Heterozygous deletion

Biological Impact

The SMAD family member 4 (SMAD4) gene encodes a transcription factor that acts as a downstream effector in the TGF- β signaling pathway. Upon phosphorylated and activated by serine-threonine receptor kinase, Smad4 is the Co-Smad which recruits other activated R-Smad proteins to the Smad transcriptional complex and regulate TGF- β -targeted genes^[72]. Smad4 has been identified as a haploinsufficient gene with one copy loss may lead to a weak protein expression and is insufficient to execute its original physiological function^[73]. SMAD4 germline mutations are associated with juvenile polyposis syndrome (JPS)^{[74][75][76][77]}. Somatic mutations of SMAD4 are commonly observed in pancreatic cancer^[78], colorectal cancer (CRC)^{[76][79][80]}, and less frequently seen in other cancers such as lung adenocarcinoma^[81],





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head and neck cancer^{[82][83]}, and cutaneous squamous cell carcinoma^[84].

Therapeutic and prognostic relevance

In Chinese patients with metastatic colorectal cancer, SMAD4 or NF1 mutations are suggested as a potential biomarker for poor prognosis to cetuximab-based therapy^[85]. Preclinical data demonstrated that depletion of SMAD4 by shRNA knockdown increased clonogenic survival and cetuximab resistance in HPV-negative head and neck squamous cell carcinoma cells^[86].

SMAD4 is also suggested as a predictive marker for 5-fluorouracil-based chemotherapy in colorectal cancer (CRC)^{[87][88]}. CRC patients with normal SMAD4 diploidy exhibited three-fold higher benefit of 5-FU/mitomycin-based adjuvant therapy when compared with those with SMAD4 deletion^[89].

Results from clinical and meta-analyses showed that loss of SMAD4 in CRC, pancreatic cancer was correlated with poor prognosis^{[90][91][92][93][94][95][96][97]}. In cervical cancer patients, weak cytoplasmic SMAD4 expression and absent nuclear SMAD4 expression were shown to be significantly associated with poor disease-free and overall 5-year survival^[98].

STK11 Heterozygous deletion

Biological Impact

The serine/threonine kinase 11 (STK11, also known as LKB1) gene encodes the multifunctional serine/threonine kinase, a tumor suppressor that functions as an inhibitor for the mTOR signaling pathway^{[99][100]}. STK11 is a haploinsufficient gene with one copy loss may lead to weak protein expression and is insufficient to execute its original physiological functions^{[101][102]}. In the mouse model, loss of STK11 promotes aggressive endometrial and squamous cell carcinomas^{[103][104]}. Mutations in STK11 have been found in lung, breast, cervical, testicular, and liver cancers, as well as malignant melanoma, pancreatic and biliary carcinoma^[105]. Germline mutations in STK11 are found in 30-70% of Peutz-Jeghers syndrome^[106].

Therapeutic and prognostic relevance

A clinical study in a pancreatic cancer patient with Peutz-Jeghers syndrome whose tumor harboring an STK11 D194E mutation coupled with the loss of heterozygosity of the other STK11 allele displayed partial response to the everolimus treatment^[107]. In another clinical case study, an adrenocorticotropic pituitary carcinoma patient whose tumor bearing an STK11 inactivating mutation responded to a combination of everolimus and radiotherapy^[108].

Preclinical data suggested that lung cancer cell lines with STK11 inactivating mutations may confer increased sensitivity to the MEK-1 and MEK-2 inhibitor, trametinib^[109].

Inactivating mutations of STK11 was shown to be associated with resistance to immune checkpoint blockade in KRAS-mutant lung adenocarcinoma (LUAC) and NSCLC (DOI: 10.1200/JCO.2017.35.15_suppl.9016)^{[110][111][112]}. It was proposed that loss of STK11 negatively impacts the number and function of tumor-infiltrating T cells (TILs) and PD-L1 expression on tumor cells and therefore results in an ineffective response to PD-1-targeting antibodies^[113].





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US FDA-APPROVED DRUG(S)

Binimetinib (MEKTOVI)

Binimetinib is an oral kinase inhibitor that targets MEK. Binimetinib is developed and marketed by Array BioPharma under the trade name MEKTOVI.

- FDA Approval Summary of Binimetinib (MEKTOVI)

NECTO (1114)	Melanoma (Approved on 2018/06/27)
MEKTOVI ^[114]	BRAF V600E/K
NCT01909453	Encorafenib + binimetinib vs. Vemurafenib [PFS(M): 14.9 vs. 7.3]

Cobimetinib (COTELLIC)

Cobimetinib is a reversible inhibitor which targets MEK1 and MEK2. Cobimetinib is developed by Exelixis and Genentech, and marketed by Genentech under the trade name COTELLIC.

- FDA Approval Summary of Cobimetinib (COTELLIC)

DDIM[115]	Melanoma (Approved on 2015/11/10)
coBRIM ^[115] NCT01689519	BRAF V600E/K
NC101669519	Cobimetinib + vemurafenib vs. Placebo + vemurafenib [PFS(M): 12.3 vs. 7.2]

Erdafitinib (BALVERSA)

Erdafitinib is a kinase inhibitor that binds to and inhibits enzymatic activity of FGFR1, FGFR2, FGFR3 and FGFR4 based on in vitro data. Erdafitinib also binds to RET, CSF1R, PDGFRA, PDGFRB, FLT4, KIT, and VEGFR2. Erdafitinib is developed and marketed by Janssen under the trade name BALVERSA.

- FDA Approval Summary of Erdafitinib (BALVERSA)

Study DI C2004	Bladder urothelial carcinoma (Approved on 2019/04/12)
Study BLC2001 NCT02365597	-
	Erdafitinib [ORR(%): 32.2]

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)

RADIANT-4 ^[116] NCT01524783	Lung or gastrointestinal neuroendocrine tumor (Approved on 2016/02/26)
	-
	Everolimus vs. Placebo [PFS(M): 11 vs. 3.9]
BOLERO-2 ^[117] NCT00863655	Breast cancer (Approved on 2012/07/20)
	ER+/HER2-
	Everolimus + exemestane vs. Placebo + exemestane [PFS(M): 7.8 vs. 3.2]





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EXIST-2	Tuberous sclerosis complex (tsc)-associated renal angiomyolipoma (Approved on 2012/04/26)
NCT00790400	-
	Everolimus vs. Placebo [ORR(%): 41.8 vs. 0]
DADIANT O[118]	Pancreatic neuroendocrine tumor (Approved on 2011/05/05)
RADIANT-3 ^[118]	-
NCT00510068	Everolimus vs. Placebo [PFS(M): 11 vs. 4.6]
	Subependymal giant cell astrocytoma (Approved on 2010/10/29)
EXIST-1 ^[119] NCT00789828	- 1
	Everolimus vs. Placebo [ORR(%): 35.0]
RECORD-1 ^[120] NCT00410124	Renal cell carcinoma (Approved on 2009/05/30)
	Everolimus vs. Placebo [PFS(M): 4.9 vs. 1.9]

Infigratinib (TRUSELTIQ)

Infigratinib a kinase inhibitor. Infigratinib is developed and marketed by QED Therapeutics, Inc. under the trade name TRUSELTIQ.

- FDA Approval Summary of Infigratinib (TRUSELTIQ)

CDC 1200V2204	Cholangiocarcinoma (Approved on 2021/05/28)
CBGJ398X2204 NCT02150967	FGFR2 fusion
	Infigratinib [ORR(%): 23.0, DOR(M): 5]

Lenvatinib (LENVIMA)

Lenvatinib is a multiple kinase inhibitor against the VEGFR1, VEGFR2 and VEGFR3. Lenvatinib is marketed by Eisai Inc. under the trade name LENVIMA.

- FDA Approval Summary of Lenvatinib (LENVIMA)

	Endometrial carcinoma (Approved on 2021/07/22)
KEYNOTE-775 (Study 309) NCT03517449	Not microsatellite instability-high (MSI-H) or mismatch repair deficient (dMMR)
	Pembrolizumab + lenvatinib vs. Investigator's choice of doxorubicin or paclitaxel [PFS(M): 6.6
	vs. 3.8, OS(M): 17.4 vs. 12]
KEVNOTE 440	Endometrial carcinoma (Approved on 2019/09/17)
KEYNOTE-146	Not microsatellite instability high (MSI-H) or mismatch repair deficient (dMMR)
NCT02501096	Pembrolizumab + lenvatinib [ORR(%): 38.3, DOR(M): NR]
DEEL ECT[121]	Hepatocellular carcinoma (Approved on 2018/08/16)
REFLECT ^[121] NCT01761266	-
NC101761266	Lenvatinib vs. Sorafenib [OS(M): 13.6 vs. 12.3]
OF! FOT[122]	Renal cell carcinoma (Approved on 2016/05/13)
SELECT ^[122]	-
NCT01136733	Lenvatinib+ everolimus vs. Everolimus [PFS(M): 14.6 vs. 5.5]
SELECT ^[123] NCT01321554	Thyroid cancer (Approved on 2015/02/13)
	-
	Lenvatinib vs. Placebo [PFS(M): 18.3 vs. 3.6]





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Niraparib (ZEJULA)

Niraparib is an oral, small molecule inhibitor of the DNA repair enzyme poly (ADP-ribose) polymerase-1 and -2 (PARP-1, -2). Niraparib is developed and marketed by Tesaro under the trade name ZEJULA.

- FDA Approval Summary of Niraparib (ZEJULA)

PRIMA NCT02655016	Ovarian cancer, Fallopian tube cancer, Peritoneal carcinoma (Approved on 2020/04/29)
	Niraparib vs. Placebo [PFS (overall population)(M): 13.8 vs. 8.2]
	Ovarian cancer, Fallopian tube cancer, Peritoneal carcinoma (Approved on 2019/10/23)
QUADRA ^[124] NCT02354586	HRD-positive (defined by either a deleterious or suspected deleterious BRCA mutation, and/or genomic instability)
	Niraparib [ORR(%): 24.0, DOR(M): 8.3]
NOVA[125]	Ovarian cancer, Fallopian tube cancer, Peritoneal carcinoma (Approved on 2017/03/27)
NOVA ^[125]	
NCT01847274	Niraparib vs. Placebo [PFS (overall population)(M): 11.3 vs. 4.7]

Olaparib (LYNPARZA)

Olaparib is an oral, small molecule inhibitor of poly (ADP-ribose) polymerase-1, -2, and -3 (PARP-1, -2, -3). Olaparib is developed by KuDOS Pharmaceuticals and marketed by AstraZeneca under the trade name LYNPARZA.

- FDA Approval Summary of Olaparib (LYNPARZA)

OlympiA NCT02032823	Her2-negative high-risk early breast cancer (Approved on 2022/03/11)
	gBRCA
	Olaparib vs. Placebo [invasive disease-free survival (IDFS)(M):]
	Prostate cancer (Approved on 2020/05/19)
PROfound ^[126]	ATMm, BRCA1m, BRCA2m, BARD1m, BRIP1m, CDK12m, CHEK1m, CHEK2m, FANCLm
NCT02987543	PALB2m, RAD51Bm, RAD51Cm, RAD51Dm, RAD54Lm
	Olaparib vs. Enzalutamide or abiraterone acetate [PFS(M): 5.8 vs. 3.5]
	Ovarian cancer (Approved on 2020/05/08)
PAOLA-1 ^[127]	HRD-positive (defined by either a deleterious or suspected deleterious BRCA mutation
NCT02477644	and/or genomic instability)
	Olaparib + bevacizumab vs. Placebo + bevacizumab [PFS(M): 37.2 vs. 17.7]
POLO ^[128]	Pancreatic adenocarcinoma (Approved on 2019/12/27)
NCT02184195	Germline BRCA mutation (deleterious/suspected deleterious)
NC102104193	Olaparib vs. Placebo [ORR(%): 23.0 vs. 12.0, PFS(M): 7.4 vs. 3.8]
SOLO-1 ^[129]	Ovarian cancer, Fallopian tube cancer, Peritoneal carcinoma (Approved on 2018/12/19)
NCT01844986	Germline or somatic BRCA-mutated (gBRCAm or sBRCAm)
NC101044900	Olaparib vs. Placebo [PFS(M): NR vs. 13.8]
Ol A D[130]	Breast cancer (Approved on 2018/02/06)
OlympiAD ^[130]	Germline BRCA mutation (deleterious/suspected deleterious) HER2-negative
NCT02000622	Olaparib vs. Chemotherapy [PFS(M): 7 vs. 4.2]
SOLO-2/ENGOT-Ov21 ^[131] NCT01874353	Ovarian cancer, Fallopian tube cancer, Peritoneal carcinoma (Approved on 2017/08/17)
	gBRCA+
	Olaparib vs. Placebo [PFS(M): 19.1 vs. 5.5]





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Study19 ^[132] NCT00753545	Ovarian cancer, Fallopian tube cancer, Peritoneal carcinoma (Approved on 2017/08/17)
	-
	Olaparib vs. Placebo [PFS(M): 8.4 vs. 4.8]
Study 42 ^[133] NCT01078662	Ovarian cancer (Approved on 2014/12/19)
	Germline BRCA mutation (deleterious/suspected deleterious)
	Olaparib [ORR(%): 34.0, DOR(M): 7.9]

Pazopanib (VOTRIENT)

Pazopanib is an oral, small molecule, multi-kinase inhibitor that targets receptor tyrosine kinase including vascular endothelial growth factor receptor-1, -2, -3 (VEGFR-1, -2, -3), platelet-derived growth factor receptor- α , - β (PDGFR- α , - β), c-kit, fibroblast growth factor-1 and -3 (FGFR-1, -3), thereby inhibiting angiogenesis. Pazopanib is developed and marketed by GlaxoSmithKline under the trade name VOTRIENT.

- FDA Approval Summary of Pazopanib (VOTRIENT)

PALETTE ^[134] NCT00753688	Sarcoma (Approved on 2016/04/26)
	Pazopanib vs. Placebo [PFS(M): 4.6 vs. 1.6]
VEC405400[135]	Renal cell carcinoma (Approved on 2009/10/19)
VEG105192 ^[135] NCT00334282	
	Pazopanib vs. Placebo [PFS(M): 9.2 vs. 4.2]

Ponatinib (ICLUSIG)

Ponatinib is an oral, small molecule, multi-kinase inhibitor designed to inhibit the activity of the tyrosine kinase ABL, including the T315I mutated ABL as well. Ponatinib is developed and marketed by ARIAD under the trade name ICLUSIG.

- FDA Approval Summary of Ponatinib (ICLUSIG)

PACE ^[136] NCT01207440	Chronic phase chronic myeloid leukemia (Approved on 2014/03/12)
	-
	Ponatinib [MCyR(%): 55]
DA OF[136]	Accelerated phase chronic myeloid leukemia (Approved on 2014/03/12)
PACE ^[136]	
NCT01207440	Ponatinib [MaHR(%): 57]
DA OF[136]	Blast phase chronic myeloid leukemia (Approved on 2014/03/12)
PACE ^[136] NCT01207440	-
	Ponatinib [MaHR(%): 31]
PACE ^[136] NCT01207440	Philadelphia-positive acute lymphoblastic leukemia (Approved on 2014/03/12)
	-
	Ponatinib [MaHR(%): 41]





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Regorafenib (STIVARGA)

Regorafenib is a multi-kinase inhibitor which targets angiogenic, stromal and oncogenic receptor tyrosine kinases (RTKs). Regorafenib is developed and marketed by Bayer HealthCare Pharmaceuticals under the trade name STIVARGA.

- FDA Approval Summary of Regorafenib (STIVARGA)

RESORCE ^[137]	Hepatocellular carcinoma, Hepatocellular carcinoma (Approved on 2017/04/27)
NCT01774344	
NC101774344	Bsc vs. Placebo [OS(M): 10.6 vs. 7.8]
GRID ^[138]	Gastrointestinal stromal tumor (Approved on 2013/02/25)
NCT01271712	
NC1012/1/12	Regorafenib vs. Placebo [PFS(M): 4.8 vs. 0.9]
00DDE0T[139]	Colorectal cancer (Approved on 2012/09/27)
CORRECT ^[139]	- /
NCT01103323	Regorafenib vs. Placebo [OS(M): 6.4 vs. 5]

Rucaparib (RUBRACA)

Rucaparib is an inhibitor of the DNA repair enzyme poly (ADP-ribose) polymerase-1, -2 and -3 (PARP-1, -2, -3). Rucaparib is developed and marketed by Clovis Oncology under the trade name RUBRACA.

- FDA Approval Summary of Rucaparib (RUBRACA)

• • • • • • • • • • • • • • • • • • • •	
TRITONO	Prostate cancer (Approved on 2020/05/15)
TRITON2 NCT02952534	gBRCA+, sBRCA
NC102952554	Rucaparib [ORR(%): 44.0, DOR(M): NE]
	Ovarian cancer, Fallopian tube cancer, Peritoneal carcinoma (Approved on 2018/04/06)
ARIEL3[140]	AII HRD tBRCA
NCT01968213	Rucaparib vs. Placebo [PFS (All)(M): 10.8 vs. 5.4, PFS (HRD)(M): 13.6 vs. 5.4, PFS
	(tBRCA)(M): 16.6 vs. 5.4]
ARIEL2[141]	Ovarian cancer (Approved on 2016/12/19)
NCT01482715,	Germline and/or somatic BRCA mutation
NCT01891344	Rucaparib [ORR(%): 54.0]

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)

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





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Sunitinib (SUTENT)

Sunitinib is an oral, small molecule, multi-kinase inhibitor that targets receptor tyrosine kinase including platelet-derived growth factor receptor- α , - β (PDGFR- α , - β), vascular endothelial growth factor receptors-1, -2, -3 (VEGFR-1, -2, -3), c-kit, Fms-like tyrosine kinase-3 (FLT3), colony stimulating factor receptor type 1 (CSF-1R), and the glial cell-line derived neurotrophic factor receptor (RET), thereby inhibiting angiogenesis. Sunitinib is developed and marketed by Pfizer under the trade name SUTENT.

- FDA Approval Summary of Sunitinib (SUTENT)

[142][143][144]	Pancreatic cancer (Approved on 2011/05/20)
NCT00428597	
NC100420397	Sunitinib vs. Placebo [PFS(M): 10.2 vs. 5.4]
[145][146]	Renal cell carcinoma (Approved on 2007/02/02)
NCT00083889	
140100003009	Sunitinib vs. Ifn-α [PFS(W): 47.3 vs. 22]
[147][148][146]	Renal cell carcinoma (Approved on 2007/02/02)
NCT00077974	-
110100077974	Sunitinib [ORR(%): 34.0]
[148][146]	Renal cell carcinoma (Approved on 2007/02/02)
NCT00054886	
110100034000	Sunitinib [ORR(%): 36.5]
[149]	Gastrointestinal stromal tumor (Approved on 2006/01/26)
NCT00075218	-
140100073210	Sunitinib vs. Placebo [TTP(W): 27.3 vs. 6.4]

Talazoparib (TALZENNA)

Talazoparib is an inhibitor of poly (ADP-ribose) polymerase (PARP) enzymes, including PARP1 and PARP2. Talazoparib is developed and marketed by Pfizer under the trade name TALZENNA.

- FDA Approval Summary of Talazoparib (TALZENNA)

EMBRACA ^[150]	Breast cancer (Approved on 2018/10/16)
	Germline BRCA mutation (deleterious/suspected deleterious) HER2-negative
NCT01945775	Talazoparib vs. Chemotherapy [PFS(M): 8.6 vs. 5.6]

Temsirolimus (TORISEL)

Temsirolimus is a soluble ester of sirolimus (rapamycin, brand-name drug Rapamune) and functions as an inhibitor of mammalian target of rapamycin complex (mTORC). The inhibitory molecular mechanism is similar to Everolimus. Temsirolimus is developed by Wyeth Pharmaceuticals and marketed by Pfizer under the trade name TORISEL.

- FDA Approval Summary of Temsirolimus (TORISEL)

[151]	Renal cell carcinoma (Approved on 2007/05/30)
1 - 1	
NCT00065468	Temsirolimus vs. Ifn-α [OS(M): 10.9 vs. 7.3]





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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)

BRF117019 ^[152]	Anaplastic thyroid cancer (Approved on 2018/05/04)				
NCT02034110	BRAF V600E				
NC102034110	Dabrafenib + trametinib [ORR(%): 61.0]				
BRF113928 ^[153]	Non-small cell lung cancer (Approved on 2017/06/22)				
NCT01336634	BRAF V600E				
NC101330034	Trametinib + dabrafenib vs. Dabrafenib [ORR(%): 63.0 vs. 27.0, DOR(M): 12.6 vs. 9.9]				
OOMBL -1[154]	Melanoma (Approved on 2014/01/10)				
COMBI-d ^[154] NCT01584648	BRAF V600E/K				
NC101304040	Trametinib + dabrafenib vs. Dabrafenib + placebo [PFS(M): 9.3 vs. 8.8]				
METRIC ^[155]	Melanoma (Approved on 2013/05/29)				
NCT01245062	BRAF V600E/K				
NC101243002	Trametinib vs. Dacarbazine or paclitaxel [PFS(M): 4.8 vs. 1.5]				

D=day; W=week; M=month





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ONGOING CLINICAL TRIALS

Trials were searched by applying filters: study status, patient's diagnosis, intervention, location and/or biomarker(s). Please visit https://clinicaltrials.gov to search and view for a complete list of open available and updated matched trials.

No trial has been found.





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SUPPLEMENTARY INFORMATION OF TESTING RESULTS DETAILED INFORMATION OF VARIANTS WITH CLINICAL RELEVANCE

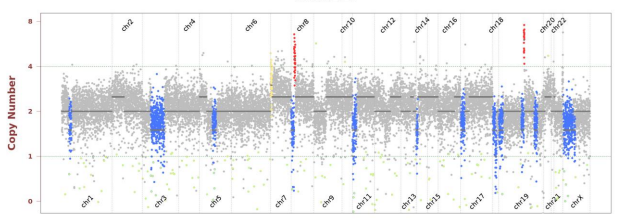
- Single Nucleotide and Small InDel Variants

Gene	Amino Acid Change	Exon cDNA Chan		Accession COSMIC		Allele Frequency	Coverage	
BRAF	V600_K601delinsE	15	c.1799_1801del	NM_004333	COSM1133	31.9%	1336	
TP53	Splice donor	_	c.993+1G>A	NM 000546	COSM44295	42.9%	638	

- Copy Number Alterations

Observed copy number (CN) for each evaluated position is shown on the y-axis. Regions referred to as amplification or deletion are shown in color. Regions without significant changes are represented in gray.

AA-22-02113









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OTHER DETECTED VARIANTS

Gene	Amino Acid Change	Exon	cDNA Change	Accession Number	COSMIC ID	Allele Frequency	Coverage
DOT1L	D588E	18	c.1764C>G	NM_032482	-	75.5%	339
EPHA2	E375*	5	c.1123G>T	NM_004431	-	6.1%	440
EPHA2	K468N	6	c.1404G>T	NM_004431	-	7.5%	346
EPHA2	W467*	6	c.1401G>A	NM_004431	-	7.5%	346
ERBB2	D483N	12	c.1447G>A	NM_004448	COSM6906406	53.5%	400
FANCL	Splice region	-	c.155+7T>C	NM_018062	-	49.4%	1158
FAT1	A3644M	19	c.10930_10931delinsAT	NM_005245	-	51.4%	1212
FAT1	Y4232C	25	c.12695A>G	NM_005245	-	54.1%	747
IRS2	R1097C	1	c.3289C>T	NM_003749	-	51.2%	168
KMT2D	G3660A	39	c.10979G>C	NM_003482	-	50.0%	178
NOTCH4	G1196R	21	c.3586G>A	NM_004557	-	44.2%	156
VEGFB	Splice region	-	c.411-3T>C	NM_003377	-	42.8%	711

Note:

- This table enlists variants detected by the panel other than those with clinical relevance (reported in Testing Result section).

The clinical impact of a genetic variant is determined according to ACT Genomics in-house clinical knowledge database. A negative result does not necessarily indicate absence of biological effect on the tumor. Some variants listed here may possibly have preclinical data or may show potential clinical relevance in the future.





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TEST DETAILS SPECIMEN RECEIVED AND PATHOLOGY REVIEW





- Collection date: Jan 2022Facility retrieved: 臺北榮總
- H&E-stained section No.: S11170160A
- Collection site: Pancreas
- Examined by: Dr. Yeh-Han Wang
 - 1. The percentage of viable tumor cells in total cells in the whole slide (%): 60%
 - 2. The percentage of viable tumor cells in total cells in the encircled areas in the whole slide (%): 60%
 - 3. The percentage of necrotic cells (including necrotic tumor cells) in total cells in the whole slide (%): 0%
 - 4. The percentage of necrotic cells (including necrotic tumor cells) in total cells in the encircled areas in the whole slide (%): 0%
 - 5. Additional comment: NA
- Manual macrodissection: Not performed
- The outline highlights the area of malignant neoplasm annotated by a pathologist.

RUN QC

Panel: ACTOnco[®]+

DNA test

- Mean Depth: 795x
- Target Base Coverage at 100x: 94%

RNA test

- Average unique RNA Start Sites per control GSP2: 164





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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.
- The possibility cannot be excluded that certain pathogenic variants detected by other sequencing tools may not be reported in the test because of technical limitation of bioinformatics algorithm or the NGS sequencing platform, e.g. low coverage.
- 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.

NEXT-GENERATION SEQUENCING (NGS) METHODS

Extracted genomic DNA was amplified using primers targeting coding exons of analyzed genes and subjected to library construction. Barcoded libraries were subsequently conjugated with sequencing beads by emulsion PCR and enriched using Ion Chef system. Sequencing was performed according to Ion Proton or Ion S5 sequencer protocol (Thermo Fisher Scientific).

Raw reads generated by the sequencer were mapped to the hg19 reference genome using the Ion Torrent Suite. 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. VEP (Variant Effect Predictor) was used to annotate every variant using databases from Clinvar, COSMIC and Genome Aggregation database. Variants with coverage ≥ 25, allele frequency ≥ 5% and actionable variants with allele frequency ≥ 2% were retained. This test provides uniform coverage of the targeted regions, enabling target base coverage at 100x ≥ 85% with a mean coverage ≥ 500x.

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

Tumor mutational burden (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 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 ≥ 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).





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RNA test

Extracted RNA was reverse-transcribed and subjected to library construction. Sequencing was performed according to Ion Proton or Ion S5 sequencer protocol (Thermo Fisher Scientific). To ensure sequencing quality for fusion variant analysis, the average unique RNA Start Sites (SS) per control Gene Specific Primer 2 (GSP 2) should be ≥ 10.

The fusion analysis pipeline aligned sequenced reads to the human reference genome, identified regions that map to noncontiguous regions of the genome, applied filters to exclude probable false-positive events and, annotated previously characterized fusion events according to Quiver Gene Fusion Database, a curated database owned and maintained by ArcherDX. In general, samples with detectable fusions need to meet the following criteria: (1) Number of unique start sites (SS) for the GSP2 ≥ 3; (2) Number of supporting reads spanning the fusion junction ≥ 5; (3) Percentage of supporting reads spanning the fusion junction ≥ 10%; (4) Fusions annotated in Quiver Gene Fusion Database.

DATABASE USED

- Reference genome: Human genome sequence hg19
- COSMIC v.92
- Genome Aggregation database r2.1.1
- ClinVar (version 20210404)
- ACT Genomics in-house database
- Quiver Gene Fusion Database version 5.1.18

Variant Analysis:

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醫檢師張筑芫 博士 Chu-Yuan Chang Ph.D. 檢字第 020115 號

Sign Off







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GENE LIST SNV & CNV

ABCB1*	ABCC2*	ABCG2*	ABL1	ABL2	ADAMTS1	ADAMTS13	ADAMTS15	ADAMTS16	ADAMTS18	ADAMTS6	ADAMTSS
ADAMTSL1	ADGRA2	ADH1C*	AKT1	AKT2	AKT3	ALDH1A1*	ALK	AMER1	APC	AR	ARAF
ARID1A	ARID1B	ARID2	ASXL1	ATM	ATR	ATRX	AURKA	AURKB	AXIN1	AXIN2	AXL
B2M	BAP1	BARD1	BCL10	BCL2*	BCL2L1	BCL2L2*	BCL6	BCL9	BCOR	BIRC2	BIRC3
BLM	BMPR1A	BRAF	BRCA1	BRCA2	BRD4	BRIP1	BTG1	BTG2*	BTK	BUB1B	CALR
CANX	CARD11	CASP8	CBFB	CBL	CCNA1	CCNA	CCNB1	CCNB2	CCNB3	CCND1	CCND2
CCND3	CCNE1	CCNE2	CCNH	CD19	CD274	CD58	CD70*	CD79A	CD79B	CDC73	CDH1
CDK1	CDK12	CDK2	CDK4	CDK5	CDK6	CDK7	CDK8	CDK9	CDKN1A	CDKN1B	CDKN2A
CDKN2B	CDKN2C	CEBPA*	CHEK1	CHEK2	CIC	CREBBP	CRKL	CRLF2	CSF1R	CTCF	CTLA4
CTNNA1	CTNNB1	CUL3	CYLD	CYP1A1*	CYP2B6*	CYP2C19*	CYP2C8*	CYP2D6	CYP2E1*	CYP3A4*	CYP3A5*
DAXX	DCUN1D1	DDR2	DICER1	DNMT3A	DOT1L	DPYD	DTX1	E2F3	EGFR	EP300	EPCAM
EPHA2	ЕРНА3	EPHA5	ЕРНА7	EPHB1	ERBB2	ERBB3	ERBB4	ERCC1	ERCC2	ERCC3	ERCC4
ERCC5	ERG	ESR1	ESR2	ETV1	ETV4	EZH2	FAM46C	FANCA	FANCC	FANCD2	FANCE
FANCF	FANCG	FANCL	FAS	FAT1	FBXW7	FCGR2B	FGF1*	FGF10	FGF14	FGF19*	FGF23
FGF3	FGF4*	FGF6	FGFR1	FGFR2	FGFR3	FGFR4	FH	FLCN	FLT1	FLT3	FLT4
FOXL2*	FOXP1	FRG1	FUBP1	GATA1	GATA2	GATA3	GNA11	GNA13	GNAQ	GNAS	GREM1
GRIN2A	GSK3B	GSTP1*	GSTT1*	HGF	HIF1A	HIST1H1C*	HIST1H1E*	HNF1A	HR	HRAS*	HSP90AA
HSP90AB1	HSPA4	HSPA5	IDH1	IDH2	IFNL3*	IGF1	IGF1R	IGF2	IKBKB	IKBKE	IKZF1
IL6	IL7R	INPP4B	INSR	IRF4	IRS1	IRS2*	JAK1	JAK2	JAK3	JUN*	KAT6A
KDM5A	KDM5C	KDM6A	KDR	KEAP1	KIT	KMT2A	кмт2С	KMT2D	KRAS	LCK	LIG1
LIG3	LMO1	LRP1B	LYN	MALT1	MAP2K1	MAP2K2	MAP2K4	MAP3K1	МАРЗК7	MAPK1	МАРК3
MAX	MCL1	MDM2	MDM4	MED12	MEF2B	MEN1	MET	MITF	MLH1	MPL	MRE11
MSH2	MSH6	MTHFR*	MTOR	MUC16	MUC4	MUC6	митүн	MYC	MYCL	MYCN	MYD88
NAT2*	NBN	NEFH	NF1	NF2	NFE2L2	NFKB1	NFKBIA	NKX2-1*	NOTCH1	NOTCH2	<i>NOTCH3</i>
NOTCH4	NPM1	NQ01*	NRAS	NSD1	NTRK1	NTRK2	NTRK3	PAK3	PALB2	PARP1	PAX5
PAX8	PBRM1	PDCD1	PDCD1LG2	PDGFRA	PDGFRB	PDIA3	PGF	PHOX2B*	PIK3C2B	PIK3C2G	РІКЗСЗ
PIK3CA	РІКЗСВ	PIK3CD	PIK3CG	PIK3R1	PIK3R2	PIK3R3	PIM1	PMS1	PMS2	POLB	POLD1
POLE	PPARG	PPP2R1A	PRDM1	PRKAR1A	PRKCA	PRKCB	PRKCG	PRKCI	PRKCQ	PRKDC	PRKN
PSMB8	PSMB9	PSME1	PSME2	PSME3	PTCH1	PTEN	PTGS2	PTPN11	PTPRD	PTPRT	RAC1
RAD50	RAD51	RAD51B	RAD51C	RAD51D	RAD52	RAD54L	RAF1	RARA	RB1	RBM10	RECQL4
REL	RET	RHOA	RICTOR	RNF43	ROS1	RPPH1	RPTOR	RUNX1	RUNX1T1	RXRA	SDHA
SDHB	SDHC	SDHD	SERPINB3	SERPINB4	SETD2	SF3B1	SGK1	SH2D1A*	SLC19A1*	SLC22A2*	SLCO1B1
SLCO1B3*	SMAD2	SMAD3	SMAD4	SMARCA4	SMARCB1	SMO	SOCS1*	SOX2*	SOX9	SPEN	SPOP
SRC	STAG2	STAT3	STK11	SUFU	SYK	SYNE1	TAF1	TAP1	TAP2	TAPBP	TBX3
TEK	TERT	TET1	TET2	TGFBR2	TMSB4X*	TNF	TNFAIP3	TNFRSF14	TNFSF11	TOP1	TP53
TPMT*	TSC1	TSC2	TSHR	TYMS	U2AF1	UBE2A*	UBE2K	UBR5	UGT1A1*	USH2A	VDR*
VEGFA	VEGFB	VHL	WT1	XIAP	XPO1	XRCC2	ZNF217				

^{*}Analysis of copy number alterations NOT available.

FUSION

		TCTD.	CCCD4		ECED2							
	BRAF	EGFR	FGFR1	FGFR2	FGFR3	MET	NRG1	NTRK1	NTRK2	NTRK3	RET	ROS1





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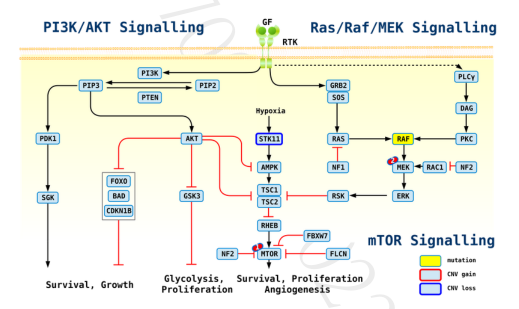
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APPENDIX

POSSIBLE THERAPEUTIC IMPLICATIONS FOR HETEROZYGOUS DELETION

Gene	Therapies	Possible effect
STK11	Binimetinib, Cobimetinib, Trametinib, Everolimus, Temsirolimus	sensitive
BAP1	Niraparib, Olaparib, Rucaparib, Talazoparib	sensitive
SMAD4	Cetuximab	resistant

SIGNALING PATHWAYS AND MOLECULAR-TARGETED AGENTS



1: Everolimus, Temsirolimus; 2: Selumetinib, Trametinib, Binimetinib, Cobimetinib



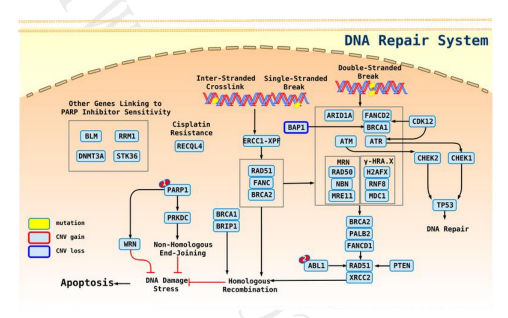


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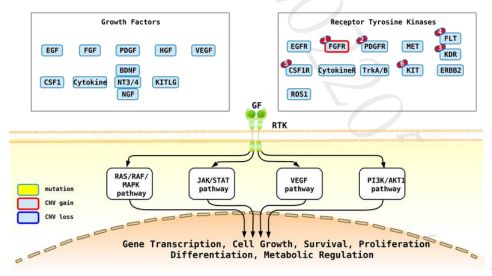
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1: Olaparib, Niraparib, Rucaparib, Talazoparib; 2: Ponatinib

Receptor Tyrosine Kinase/Growth Factor Signalling



1: Ponatinib, Lenvatinib, Erdafitinib, Infigratinib, Pazopanib; 2: Ponatinib, Pazopanib, Sunitinib, Regorafenib; 3: Ponatinib, Lenvatinib, Pazopanib, Sunitinib; 4: Lenvatinib, Pazopanib, Sunitinib; 5: Sunitinib; 6: Ponatinib, Regorafenib, Lenvatinib, Pazopanib, Sunitinib





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