

Department of Pathology and Laboratory Medicine No.201, Sec. 2, Shipai Rd., Beitou District, Taipei City, Taiwan 11217, R.O.C.

Tel: 02-2875-7449

Date: 03 Jun 2021 1 of 15

Sample Information

Patient Name: 黃書健 Gender: Male ID No.: E102374106 History No.: 46991622

Age: 77

Ordering Doctor: DOC8675J 黎洛 Ordering REQ.: 0BGKUFY **Signing in Date: 2021/06/02**

Path No.: S110-98912 MP No.: TM21009

Assay: Oncomine Tumor Mutation Load Assay

Sample Type: FFPE Block No.: S110-17521A Percentage of tumor cells: 50%

Note:

Sample Cancer Type: Liver Cancer

Table of Contents	Page
Variants (Exclude variant in Taiwan BioBank with >1% allele frequency)	2
Biomarker Descriptions	3
Relevant Therapy Summary	4
Relevant Therapy Details	5
Clinical Trials Summary	11

Report Highlights 1 Relevant Biomarkers 1 Therapies Available

17 Clinical Trials

Relevant Liver Cancer Variants

Gene	Finding
NTRK1	Not detected
NTRK3	Not detected

Relevant Biomarkers

Tier	Genomic Alteration	Relevant Therapies (In this cancer type)	Relevant Therapies (In other cancer type)	Clinical Trials
	Tumor Mutational Burden	pembrolizumab 1	pembrolizumab	17
	5.84 Mut/Mb measured			

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

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

DNA Sequence Variants

Gene	Amino Acid Change	Coding	Variant ID	Locus	Allele Frequency	Transcript	Variant Effect	Coverage
ABL2	p.(=)	c.189T>A		chr1:179100603	53.68%	NM_005158.4	synonymous	1669
MARK1	p.(R459G)	c.1375C>G		chr1:220809273	44.47%	NM_018650.4	missense	1466
MSH2	p.(I169V)	c.505A>G		chr2:47637371	44.19%	NM_000251.2	missense	1998
TCF7L1	p.(T553I)	c.1658C>T		chr2:85536476	47.14%	NM_031283.2	missense	963
AFF3	p.(E56D)	c.168A>C		chr2:100625355	16.47%	NM_001025108.1	missense	1998
LTF	p.(R23dup)	c.68_69insAAG		chr3:46501284	100.00%	NM_002343.5	nonframeshift Insertion	1973
ATR	p.(A1550V)	c.4649C>T		chr3:142231305	66.38%	NM_001184.3	missense	1294
BCL6	p.(S308L)	c.923C>T		chr3:187447270	35.34%	NM_001706.4	missense	1095
PDGFRA	p.(=)	c.1701A>G		chr4:55141055	99.94%	NM_006206.5	synonymous	1600
MYB	p.(R454C)	c.1360C>T		chr6:135518255	12.36%	NM_001130173.1	missense	1100
TNFAIP3	p.(S79N)	c.236G>A		chr6:138192600	53.14%	NM_001270507.1	missense	1993
SYNE1	p.(=)	c.25146C>T		chr6:152462438	48.82%	NM_182961.3	synonymous	1477
AKAP9	p.(S2777G)	c.8329A>G		chr7:91712652	6.70%	NM_005751.4	missense	2000
TAF1L	p.(=)	c.462G>A		chr9:32635116	5.05%	NM_153809.2	synonymous	99
FANCG	p.(?)	c8C>T		chr9:35079529	47.39%	NM_004629.1	unknown	1321
RALGDS	p.(=)	c.2646G>T		chr9:135974070	6.78%	NM_001271775.1	synonymous	59
RET	p.(V292M)	c.874G>A		chr10:43601830	63.33%	NM_020975.4	missense	1268
КАТ6В	p.(E1100fs)	c.3298_3299delGA		chr10:76781913	8.90%	NM_012330.3	frameshift Deletion	1045
KAT6B	p.(E1103fs)	c.3307_3308delGA		chr10:76781922	89.95%	NM_012330.3	frameshift Deletion	1045
CYP2C19	p.(R186C)	c.556C>T		chr10:96540330	30.03%	NM_000769.2	missense	1995
FGFR2	p.(S372C)	c.1115C>G		chr10:123274803	42.44%	NM_000141.4	missense	1998
KMT2D	p.(L915V)	c.2743C>G		chr12:49444723	47.82%	NM_003482.3	missense	1491
EP400	p.(=)	c.8250G>A		chr12:132547162	100.00%	NM_015409.4	synonymous	438
BIVM- ERCC5	p.(I746M)	c.2238A>G		chr13:103514060	51.59%	NM_001204425.1	missense	1702
ERCC5	p.(I292M)	c.876A>G		chr13:103514060	51.59%	NM_000123.3	missense	1702
PML	p.(A601D)	c.1802C>A		chr15:74335421	10.11%	NM_033238.2	missense	1999
CDH5	p.(I517T)	c.1550_1551delTCins CT		chr16:66432423	54.17%	NM_001795.4	missense	1979
NF1	p.(=)	c.2463T>C		chr17:29556096	3.70%	NM_001042492.2	synonymous	54
SEPT9	p.(=)	c.1446G>A		chr17:75488768	54.80%	NM_001113491.1	synonymous	2000
BCL2	p.(A42fs)	c.124delG		chr18:60985775	67.76%	NM_000633.2	frameshift Deletion	245

Date: 03 Jun 2021 3 of 15

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

DNA Sequence Variants (continued)

Gene	Amino Acid Change	Coding	Variant ID	Locus	Allele Frequency	Transcript	Variant Effect	Coverage
BCL2	p.(P40fs)	c.119_127delCGGGG GCCGinsGGGGCCA		chr18:60985773	32.24%	NM_000633.2	frameshift Block Substitution	245
AXL	p.(S250R)	c.750C>A		chr19:41737170	60.26%	NM_021913.4	missense	1744

Biomarker Descriptions

Tumor Mutational Burden

Background: Tumor mutational burden (TMB), also known as tumor mutational load (TML), is the count of somatic mutations in the DNA of cancer cells. TMB is determined by next-generation sequencing and is expressed as the number of mutations per megabase (mut/Mb) of DNA coding sequence¹. Errors in DNA repair, including mutations in the POLE gene and in mismatch repair (MMR) genes, are associated with increased TMB^{2,3,4,5,6}. High TMB is associated with increased neo-antigen burden and has been linked to response to immune checkpoint inhibitors (ICIs) that target the cytotoxic T lymphocyte antigen-4 (CTLA4), programmed death protein 1 (PD1), and programmed death-ligand 1 (PD-L1) inhibitors^{7,8,9,10}.

Alterations and prevalence: In one study of over 100,000 tumor samples, the median TMB value was 3.6 mut/Mb although TMB values vary widely across cancers¹¹. Certain childhood cancers, leukemia, glioblastoma, and neuroblastoma typically have low mutation burden and median TMB values <1 mut/Mb^{8,11}. In comparison, cancers that experience genotoxic insults including skin cancer and lung cancer have higher median TMB values of approximately 10 mut/Mb^{8,11}. For example, within non-small cell lung cancer (NSCLC), higher TMB was observed in former/current smokers (10.5 mut/Mb) relative to never smokers (0.6 mut/Mb)^{8,11,12}. There is no consensus around the definition of high and low TMB that could be applied universally to all tumor types, instead multiple sources suggest that TMB status is a cancer type specific attribute^{11,13,14}. In NSCLC, several studies have suggested establishing a threshold between low and high TMB of 10 +/- 1 mut/Mb^{15,16,17,18}.

Potential relevance: ICIs stimulate a patient's own T-cells to kill tumors and have exhibited benefits in some patients. The first ICI to be approved by the FDA was ipilimumab (2011), an anti-CTLA4 antibody indicated for the treatment of metastatic melanoma. In 2014, anti-PD-1 antibodies, nivolumab (2014) and pembrolizumab (2014), were subsequently approved for the treatment of metastatic melanoma. Pembrolizumab was also approved (2014) for advanced esophageal squamous cell carcinoma. In 2020, the indication for pembrolizumab19 was expanded to include TMB-H (>/= 10 mut/Mb) solid tumors that have progressed on prior therapy. Indications have been expanded for these ICIs to include several other cancer types including NSCLC, advanced renal cell carcinoma, classical Hodgkin lymphoma, recurrent or metastatic squamous cell carcinoma of the head and neck, urothelial carcinoma, microsatellite instability (MSI)-High or mismatch repair deficient (dMMR) colorectal cancer, and hepatocellular carcinoma. Atezolizumab (2016), avelumab (2017), and durvalumab (2017), that target programmed death-ligand 1 (PD-L1), were subsequently approved by the FDA. However, the predictive biomarkers that underlie the clinical benefits of these approved immunotherapies, including TMB, are under active investigation. Several published studies including the CheckMate 586 and CheckMate 817 clinical trials have concluded that high TMB was associated with improved response to FDA approved checkpoint inhibitors 16,20,21. In contrast, several promising previous trials failed to show an improvement in survival outcomes between high and low TMB including CheckMate 227 (ipilimumab + nivolumab vs. chemotherapy), CheckMate 026 (nivolumab vs. chemotherapy), KEYNOTE 189 (pembrolizumab vs. chemotherapy), KEYNOTE 021 (pembrolizumab vs. pembrolizumab + chemotherapy), and Lung-MAP (nivolumab + ipilimumab vs. nivolumab). In response, suggestions to combine TMB score with PD-L1 expression as a way to increase the predictive power for patient stratification have been reported²². Nivolumab alone or in combination with ipilimumab is recommended for use in NSCLC with evidence of high TMB²³. Pembrolizumab is indicated for use in various cancer types with evidence of metastasis including Ewing sarcoma, salivary gland neoplasms, cervical cancer, uterine sarcoma, endometrial carcinoma, thyroid cancer, ovarian cancer, esophageal cancer, esophagogastric junction cancer, breast cancer, and germ cell tumors with high TMB^{24,25,26,27,28,29,30,31,32}. TMB score estimation is affected by the utilized assays, therefore efforts are underway to develop a standardized approach for score calculation with the aim to support consistent reporting of TMB values across laboratories^{33,34,35,36}.

Relevant Therapy Summary

■ In this cancer type
O In other cancer type
O In this cancer type and other cancer types
X No evidence

Relevant Therapy	FDA	NCCN	EMA	ESMO	Clinical Trials*
pembrolizumab		•	×	×	(II)
atezolizumab	×	×	×	×	(II)
atezolizumab + chemotherapy	×	×	×	×	(II)
atezolizumab, nivolumab, ipilimumab	×	×	×	×	(II)
durvalumab, tremelimumab	×	×	×	×	(II)
ipilimumab + nivolumab	×	×	×	×	(II)
ipilimumab, nivolumab	×	×	×	×	(II)
nivolumab	×	×	×	×	(II)
pembrolizumab, ipilimumab + nivolumab	×	×	×	×	(II)
chemotherapy, tremelimumab, durvalumab	×	×	×	×	(1/11)
entinostat, nivolumab	×	×	×	×	(1/11)
BAY1905254	×	×	×	×	(l)
nivolumab, ipilimumab	×	×	×	×	(l)
pembrolizumab, targinine	×	×	×	×	(1)

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

Date: 03 Jun 2021 5 of 15

Relevant Therapy Details

Current FDA Information

In this cancer type

O In other cancer type

In this cancer type and other cancer types

FDA information is current as of 2021-04-14. For the most up-to-date information, search www.fda.gov.

Tumor Mutational Burden

pembrolizumab

Cancer type: Solid Tumor Label as of: 2021-03-22 Variant class: Tumor Mutational Burden

Indications and usage:

KEYTRUDA® is a programmed death receptor-1 (PD-1)-blocking antibody indicated:

Melanoma

- for the treatment of patients with unresectable or metastatic melanoma.
- for the adjuvant treatment of patients with melanoma with involvement of lymph node(s) following complete resection.

Non-Small Cell Lung Cancer (NSCLC)

- in combination with pemetrexed and platinum chemotherapy, as first-line treatment of patients with metastatic nonsquamous NSCLC, with no EGFR or ALK genomic tumor aberrations.
- in combination with carboplatin and either paclitaxel or paclitaxel protein-bound, as first-line treatment of patients with metastatic squamous NSCLC.
- as a single agent for the first-line treatment of patients with NSCLC expressing PD-L1 [Tumor Proportion Score (TPS) ≥ 1%] as determined by an FDA-approved test, with no EGFR or ALK genomic tumor aberrations, and is:
 - stage III where patients are not candidates for surgical resection or definitive chemoradiation, or
 - metastatic.
- as a single agent for the treatment of patients with metastatic NSCLC whose tumors express PD-L1 (TPS ≥ 1%) as determined by an FDA-approved test, with disease progression on or after platinum-containing chemotherapy. Patients with EGFR or ALK genomic tumor aberrations should have disease progression on FDA-approved therapy for these aberrations prior to receiving KEYTRUDA®.

Small Cell Lung Cancer (SCLC)

• for the treatment of patients with metastatic SCLC with disease progression on or after platinum-based chemotherapy and at least one other prior line of therapy.¹

Head and Neck Squamous Cell Cancer (HNSCC)

- in combination with platinum and FU for the first-line treatment of patients with metastatic or with unresectable, recurrent HNSCC.
- as a single agent for the first-line treatment of patients with metastatic or with unresectable, recurrent HNSCC whose tumors express PD-L1 [Combined Positive Score (CPS) ≥ 1] as determined by an FDA-approved test.
- as a single agent for the treatment of patients with recurrent or metastatic HNSCC with disease progression on or after platinum-containing chemotherapy.

Classical Hodgkin Lymphoma (cHL)

- for the treatment of adult patients with relapsed or refractory cHL.
- for the treatment of pediatric patients with refractory cHL, or cHL that has relapsed after 2 or more lines of therapy.

Primary Mediastinal Large B-Cell Lymphoma (PMBCL)

- for the treatment of adult and pediatric patients with refractory PMBCL, or who have relapsed after 2 or more prior lines of therapy.
- Limitations of Use: KEYTRUDA® is not recommended for treatment of patients with PMBCL who require urgent cytoreductive therapy.

Urothelial Carcinoma

Date: 03 Jun 2021

Tumor Mutational Burden (continued)

- for the treatment of patients with locally advanced or metastatic urothelial carcinoma who are not eligible for cisplatin-containing chemotherapy and whose tumors express PD-L1 [Combined Positive Score (CPS) ≥ 10] as determined by an FDA-approved test, or in patients who are not eligible for any platinum-containing chemotherapy regardless of PD-L1 status.¹
- for the treatment of patients with locally advanced or metastatic urothelial carcinoma who have disease progression during or following platinum-containing chemotherapy or within 12 months of neoadjuvant or adjuvant treatment with platinumcontaining chemotherapy.
- for the treatment of patients with Bacillus Calmette-Guerin (BCG)-unresponsive, high-risk, non-muscle invasive bladder cancer (NMIBC) with carcinoma in situ (CIS) with or without papillary tumors who are ineligible for or have elected not to undergo cystectomy.

Microsatellite Instability-High or Mismatch Repair Deficient Cancer

- for the treatment of adult and pediatric patients with unresectable or metastatic, microsatellite instability-high (MSI-H) or mismatch repair deficient (dMMR)
 - solid tumors that have progressed following prior treatment and who have no satisfactory alternative treatment options,¹ or
 - colorectal cancer that has progressed following treatment with a fluoropyrimidine, oxaliplatin, and irinotecan.
- Limitations of Use: The safety and effectiveness of KEYTRUDA® in pediatric patients with MSI-H central nervous system cancers have not been established.

Microsatellite Instability-High or Mismatch Repair Deficient Colorectal Cancer (CRC)

• for the first-line treatment of patients with unresectable or metastatic MSI-H or dMMR colorectal cancer (CRC).

Gastric Cancer

for the treatment of patients with recurrent locally advanced or metastatic gastric or gastroesophageal junction adenocarcinoma whose tumors express PD-L1 [Combined Positive Score (CPS) ≥ 1] as determined by an FDA-approved test, with disease progression on or after 2 or more prior lines of therapy including fluoropyrimidine- and platinum-containing chemotherapy and if appropriate, HER2/neu-targeted therapy.¹

Esophageal Cancer

- for the treatment of patients with locally advanced or metastatic esophageal or gastroesophageal junction (GEJ) (tumors with epicenter 1 to 5 centimeters above the GEJ) carcinoma that is not amenable to surgical resection or definitive chemoradiation either:
 - in combination with platinum- and fluoropyrimidine-based chemotherapy, or
 - as a single agent after one or more prior lines of systemic therapy for patients with tumors of squamous cell histology that express PD-L1 (CPS ≥10) as determined by an FDA-approved test.

Cervical Cancer

for the treatment of patients with recurrent or metastatic cervical cancer with disease progression on or after chemotherapy whose tumors express PD-L1 [Combined Positive Score (CPS) ≥ 1] as determined by an FDA-approved test.¹

Hepatocellular Carcinoma (HCC)

for the treatment of patients with HCC who have been previously treated with sorafenib.

Merkel Cell Carcinoma (MCC)

■ for the treatment of adult and pediatric patients with recurrent locally advanced or metastatic Merkel cell carcinoma.¹

Renal Cell Carcinoma (RCC)

• in combination with axitinib, for the first-line treatment of patients with advanced RCC.

Endometrial Carcinoma

in combination with lenvatinib, for the treatment of patients with advanced endometrial carcinoma that is not MSI-H or dMMR, who have disease progression following prior systemic therapy and are not candidates for curative surgery or radiation.¹

Tumor Mutational Burden-High (TMB-H) Cancer

- for the treatment of adult and pediatric patients with unresectable or metastatic tumor mutational burden-high (TMB-H) [≥ 10 mutations/megabase (mut/Mb)] solid tumors, as determined by an FDA-approved test, that have progressed following prior treatment and who have no satisfactory alternative treatment options.¹
- Limitations of Use: The safety and effectiveness of KEYTRUDA® in pediatric patients with TMB-H central nervous system cancers have not been established.

Cutaneous Squamous Cell Carcinoma (cSCC)

 for the treatment of patients with recurrent or metastatic cutaneous squamous cell carcinoma that is not curable by surgery or radiation.

Triple-Negative Breast Cancer (TNBC)

• in combination with chemotherapy, for the treatment of patients with locally recurrent unresectable or metastatic TNBC whose tumors express PD-L1 [Combined Positive Score (CPS) ≥10] as determined by an FDA approved test.²

Adult Indications: Additional Dosing Regimen of 400 mg Every 6 Weeks

for use at an additional recommended dosage of 400 mg every 6 weeks for all approved adult indications.3

¹This indication is approved under accelerated approval based on tumor response rate and durability of response. Continued approval for this indication may be contingent upon verification and description of clinical benefit in the confirmatory trials.

²This indication is approved under accelerated approval based on progression-free survival. Continued approval for this indication may be contingent upon verification and description of clinical benefit in the confirmatory trials.

³This indication is approved under accelerated approval based on pharmacokinetic data, the relationship of exposure to efficacy, and the relationship of exposure to safety. Continued approval for this dosing may be contingent upon verification and description of clinical benefit in the confirmatory trials.

Reference:

https://www.accessdata.fda.gov/drugsatfda_docs/label/2021/125514s096lbl.pdf

Date: 03 Jun 2021 8 of 15

Current NCCN Information

NCCN information is current as of 2021-04-01. For the most up-to-date information, search www.nccn.org. For NCCN International Adaptations & Translations, search www.nccn.org/global/international_adaptations.aspx.

Tumor Mutational Burden

pembrolizumab

Cancer type: Cholangiocarcinoma, Liver Cancer Variant class: Tumor Mutational Burden

NCCN Recommendation category: 2A

Population segment (Line of therapy):

Gallbladder Cancer, Intrahepatic, Extrahepatic; Progression (Subsequent therapy); Useful in certain circumstances

Reference: NCCN Guidelines® - NCCN-Hepatobiliary Cancers [Version 1.2021]

O pembrolizumab

Cancer type: Chondrosarcoma, Osteosarcoma Variant class: Tumor Mutational Burden

NCCN Recommendation category: 2A

Population segment (Line of therapy):

Unresectable, Metastatic, Progression (Subsequent therapy); Useful in certain circumstances

Reference: NCCN Guidelines® - NCCN-Bone Cancer [Version 1.2021]

O pembrolizumab

Cancer type: Breast Cancer Variant class: Tumor Mutational Burden

NCCN Recommendation category: 2A

Population segment (Line of therapy):

Stage IV; Recurrent, Invasive, Unresectable, Local (Line of therapy not specified); Useful in certain circumstances

Reference: NCCN Guidelines® - NCCN-Breast Cancer [Version 3.2021]

O pembrolizumab

Cancer type: Cervical Cancer Variant class: Tumor Mutational Burden

NCCN Recommendation category: 2A

Population segment (Line of therapy):

Recurrent, Metastatic, Progression (Second-line therapy); Useful in certain circumstances

Reference: NCCN Guidelines® - NCCN-Cervical Cancer [Version 1.2021]

O pembrolizumab

Cancer type: Esophageal Cancer, Variant class: Tumor Mutational Burden

Gastroesophageal Junction Adenocarcinoma

NCCN Recommendation category: 2A

Population segment (Line of therapy):

 Adenocarcinoma, Squamous Cell; Unresectable, Locally Advanced, Recurrent, Metastatic (Second-line therapy, Subsequent therapy); Useful in certain circumstances

Reference: NCCN Guidelines® - NCCN-Esophageal and Esophagogastric Junction Cancers [Version 1.2021]

O pembrolizumab

Cancer type: Gastric Cancer Variant class: Tumor Mutational Burden

NCCN Recommendation category: 2A

Population segment (Line of therapy):

 Unresectable, Locally Advanced, Recurrent, Metastatic (Second-line therapy, Subsequent therapy); Useful in certain circumstances

Reference: NCCN Guidelines® - NCCN-Gastric Cancer [Version 2.2021]

O pembrolizumab

Cancer type: Head and Neck Cancer Variant class: Tumor Mutational Burden

NCCN Recommendation category: 2A

Population segment (Line of therapy):

- Nasopharyngeal; Recurrent, Unresectable, Metastatic (Subsequent therapy); Useful in certain circumstances
- Salivary Gland Neoplasm; Recurrent, Unresectable, Metastatic (Line of therapy not specified); Useful in certain circumstances

Reference: NCCN Guidelines® - NCCN-Head and Neck Cancers [Version 2.2021]

O pembrolizumab

Cancer type: Ovarian Cancer Variant class: Tumor Mutational Burden

NCCN Recommendation category: 2A

Population segment (Line of therapy):

■ Epithelial, Less Common Ovarian Cancers, Fallopian Tube, Primary Peritoneal; Recurrent (Recurrence therapy); Useful in certain circumstances

Reference: NCCN Guidelines® - NCCN-Ovarian Cancer [Version 1.2021]

O pembrolizumab

Cancer type: Testicular Cancer Variant class: Tumor Mutational Burden

NCCN Recommendation category: 2A

Population segment (Line of therapy):

■ Germ Cell Tumor; Metastatic (Third-line therapy); Useful in certain circumstances

Reference: NCCN Guidelines® - NCCN-Testicular Cancer [Version 1.2021]

O pembrolizumab

Cancer type: Thyroid Gland Follicular Carcinoma, Variant class: Tumor Mutational Burden

Thyroid Gland Hurthle Cell Carcinoma, Thyroid

Gland Papillary Carcinoma

NCCN Recommendation category: 2A

Population segment (Line of therapy):

Locally Recurrent, Advanced, Metastatic (Line of therapy not specified)

Reference: NCCN Guidelines® - NCCN-Thyroid Carcinoma [Version 3.2020]

O pembrolizumab

Cancer type: Thyroid Gland Medullary Carcinoma Variant class: Tumor Mutational Burden

NCCN Recommendation category: 2A

Population segment (Line of therapy):

Recurrent, Persistent, Local, Distant Metastases (Line of therapy not specified); Useful in certain circumstances

Reference: NCCN Guidelines® - NCCN-Thyroid Carcinoma [Version 3.2020]

O pembrolizumab

NCCN Recommendation category: 2A

Population segment (Line of therapy):

Stage IVC; Metastatic (Line of therapy not specified); Useful in certain circumstances

Reference: NCCN Guidelines® - NCCN-Thyroid Carcinoma [Version 3.2020]

pembrolizumab

Cancer type: Endometrial Cancer, Uterine Sarcoma Variant class: Tumor Mutational Burden

NCCN Recommendation category: 2A

Population segment (Line of therapy):

Recurrent, Metastatic (Line of therapy not specified); Useful in certain circumstances

Reference: NCCN Guidelines® - NCCN-Uterine Neoplasms [Version 1.2021]

O pembrolizumab

Cancer type: Ewing Sarcoma Variant class: Tumor Mutational Burden

NCCN Recommendation category: 2B

Population segment (Line of therapy):

Metastatic (Line of therapy not specified)

Reference: NCCN Guidelines® - NCCN-Bone Cancer [Version 1.2021]

Clinical Trials Summary

Tumor Mutational Burden

NCT ID	Title	Phase
NCT03767075	Basket of Baskets: A Modular, Open-label, Phase II, Multicentre Study To Evaluate Targeted Agents in Molecularly Selected Populations With Advanced Solid Tumours	II
NCT04185831	MEGALIT - a Multicenter, Basket and Umbrella Explorative Trial on the Efficacy and Safety of Molecular Profile Selected Commercially Available Targeted Anti-cancer Drugs in Patients With Advanced Cancers Progressive on Standard Therapy	II
NCT04589845	Tumor-Agnostic Precision Immunooncology and Somatic Targeting Rational for You (TAPISTRY) Phase II Platform Trial	II
NCT04632992	MyTACTIC: An Open-Label Phase II Study Evaluating Targeted Therapies in Patients Who Have Advanced Solid Tumors With Genomic Alterations or Protein Expression Patterns Predictive of Response	II
NCT04591431	The Rome Trial From Histology to Target: the Road to Personalize Target Therapy and Immunotherapy	II
NCT02029001	A Two-period, Multicenter, Randomized, Open-label, Phase II Study Evaluating the Clinical Benefit of a Maintenance Treatment Targeting Tumor Molecular Alterations in Patients With Progressive Locally-advanced or Metastatic Solid Tumors MOST: My own specific treatment	II
NCT03297606	Canadian Profiling and Targeted Agent Utilization Trial (CAPTUR): A Phase II Basket Trial	II
NCT03668119	A Randomized, Open-Label, Phase II Study of Nivolumab in Combination With Ipilimumab or Nivolumab Monotherapy in Participants With Advanced or Metastatic Solid Tumors of High Tumor Mutational Burden (TMB-H)	II
No NCT ID	A Multicenter Phase II Study Of Nivolumab Monotherapy In Recurrent And/Or Metastatic Gastrointestinal Cancer Patients With High Tumor Mutation Burden (TMB-H)	II
No NCT ID	TR Accompanied with the Study Titled A Multicenter Phase II Study of Nivolumab Monotherapy in Recurrent and/or Metastatic Gastrointestinal Cancer Patients with High Tumor Mutation Burden (TMB-H).	II
NCT02628067	A Clinical Trial of Pembrolizumab (MK-3475) Evaluating Predictive Biomarkers in Subjects With Advanced Solid Tumors (KEYNOTE 158)	II
NCT02693535	Targeted Agent and Profiling Utilization Registry (TAPUR) Study	II
NCT03518606	A Phase I/II Basket Trial Evaluating A Combination Of Metronomic Oral Vinorelbine Plus Anti-PD-L1/ Anti-CTLA4 ImmunothErapy In Patients With Advanced Solid Tumour	1/11
NCT03838042	INFORM2 Exploratory Multinational Phase I/II Combination Study of Nivolumab and Entinostat in Children and Adolescents with Refractory High-risk Malignancies	1/11

12 of 15

Clinical Trials Summary (continued)

Tumor Mutational Burden (continued)

NCT ID	Title	Phase
NCT03666273	An Open-label, Phase I, First-in-human, Dose Escalation and Expansion Study to Evaluate the Safety, Tolerability, Maximum Tolerated or Administered Dose, Pharmacokinetics, Pharmacodynamics and Tumor Response Profile of the ILDR2 Function-blocking Antibody BAY1905254 in Patients With Advanced Solid Tumors	I
NCT04500548	3CI Study: Childhood Cancer Combination Immunotherapy. Phase Ib and Expansion Study of Nivolumab Combination Immunotherapy in Children, Adolescent, and Young Adult (CAYA) Patients With Relapsed/Refractory Hypermutant Cancers	I
NCT03236935	Phase Ib Trial of L-NMMA in Combination With Pembrolizumab in Patients With Melanoma, Non-Small Cell Lung Cancer, Head and Neck Squamous Cell Carcinoma, Classical Hodgkin Lymphoma, Urothelial Carcinoma, Cervical Cancer, Esophageal Cancer, Gastric Cancer, Hepatocellular Carcinoma, Merkel Cell Carcinoma, Primary Mediastinal Large B-cell Lymphoma, Renal Cell Carcinoma, Small Cell Lung Cancer, Microsatellite Instability-High/Mismatch Repair Deficient Cancer, or for the Treatment of Adult Patients With Unresectable or Metastatic Tumor Mutational Burden-High Solid Tumors	I

Date: 03 Jun 2021 13 of 15

Signatures

Testing Personnel:

Laboratory Supervisor:

Pathologist:

Date: 03 Jun 2021

References

- Yuan et al. Novel technologies and emerging biomarkers for personalized cancer immunotherapy. 4:3. PMID: 26788324
- Temko et al. Somatic POLE exonuclease domain mutations are early events in sporadic endometrial and colorectal carcinogenesis, determining driver mutational landscape, clonal neoantigen burden and immune response. J. Pathol. 2018 Jul;245(3):283-296. PMID: 29604063
- 3. Mehnert et al. Immune activation and response to pembrolizumab in POLE-mutant endometrial cancer. 126(6):2334-40. PMID: 27159395
- 4. Bouffet et al. Immune Checkpoint Inhibition for Hypermutant Glioblastoma Multiforme Resulting From Germline Biallelic Mismatch Repair Deficiency. J. Clin. Oncol. 2016 Jul 1;34(19):2206-11. PMID: 27001570
- 5. Humphris et al. Hypermutation In Pancreatic Cancer, Gastroenterology, 2017 Jan;152(1):68-74.e2. PMID: 27856273
- 6. Bonneville et al. Landscape of Microsatellite Instability Across 39 Cancer Types. JCO Precis Oncol. 2017;2017. PMID: 29850653
- 7. Snyder et al. Genetic basis for clinical response to CTLA-4 blockade in melanoma. N. Engl. J. Med. 2014 Dec 4;371(23):2189-2199. PMID: 25409260
- 8. Alexandrov et al. Signatures of mutational processes in human cancer. Nature. 2013 Aug 22;500(7463):415-21. PMID: 23945592
- Rizvi et al. Mutational landscape determines sensitivity to PD-1 blockade in non-small cell lung cancer. Science. 2015 Apr 3;348(6230):124-8. PMID: 25765070
- 10. Van et al. Genomic correlates of response to CTLA-4 blockade in metastatic melanoma. Science. 2015 Oct 9;350(6257):207-211. PMID: 26359337
- 11. Chalmers et al. Analysis of 100,000 human cancer genomes reveals the landscape of tumor mutational burden. 9:34. PMID: 28420421
- 12. Govindan et al. Genomic landscape of non-small cell lung cancer in smokers and never-smokers. Cell. 2012 Sep 14;150(6):1121-34. PMID:22980976
- 13. Endris et al. Measurement of tumor mutational burden (TMB) in routine molecular diagnostics: in silico and real-life analysis of three larger gene panels. Int. J. Cancer. 2019 May 1;144(9):2303-2312. PMID: 30446996
- 14. Meléndez et al. Methods of measurement for tumor mutational burden in tumor tissue. Transl Lung Cancer Res. 2018 Dec;7(6):661-667. PMID: 30505710
- 15. Hellmann et al. Nivolumab plus Ipilimumab in Lung Cancer with a High Tumor Mutational Burden. N. Engl. J. Med. 2018 May 31;378(22):2093-2104. PMID: 29658845
- 16. Ready et al. First-Line Nivolumab Plus Ipilimumab in Advanced Non-Small-Cell Lung Cancer (CheckMate 568): Outcomes by Programmed Death Ligand 1 and Tumor Mutational Burden as Biomarkers. J. Clin. Oncol. 2019 Apr 20;37(12):992-1000. PMID: 30785829
- 17. Alborelli et al. Tumor mutational burden assessed by targeted NGS predicts clinical benefit from immune checkpoint inhibitors in non-small cell lung cancer. J. Pathol. 2020 Jan;250(1):19-29. PMID: 31471895
- 18. Heeke et al. In-house Implementation of Tumor Mutational Burden Testing to Predict Durable Clinical Benefit in Non-small Cell Lung Cancer and Melanoma Patients. Cancers (Basel). 2019 Aug 29;11(9). PMID: 31470674
- 19. https://www.accessdata.fda.gov/drugsatfda_docs/label/2021/125514s096lbl.pdf
- 20. F. et al. CheckMate 817: First-Line Nivolumab + Ipilimumab in Patients with ECOG PS 2 and Other Special Populations with Advanced NSCLC. Volume 14, Issue 10, Supplement, Pages S214–S215. Abstract OA04.02
- 21. Zhu et al. Association Between Tumor Mutation Burden (TMB) and Outcomes of Cancer Patients Treated With PD-1/PD-L1 Inhibitions: A Meta-Analysis. Front Oncol, 9:1161, 04 Nov 2019. PMID: 31258479
- 22. Castellanos et al. Tumor mutational burden (TMB) and PD-L1 expression as predictors of response to immunotherapy (IO) in NSCLC. Journal of Clinical Oncology 37, no. 15_suppl (May 20, 2019) 2630-2630. Abstract 2630
- 23. NCCN Guidelines® NCCN-Non-Small Cell Lung Cancer [Version 4.2021]
- 24. NCCN Guidelines® NCCN-Bone Cancer [Version 1.2021]
- 25. NCCN Guidelines® NCCN-Head and Neck Cancers [Version 2.2021]
- 26. NCCN Guidelines® NCCN-Testicular Cancer [Version 1.2021]
- 27. NCCN Guidelines® NCCN-Cervical Cancer [Version 1.2021]
- 28. NCCN Guidelines® NCCN-Uterine Neoplasms [Version 1.2021]
- 29. NCCN Guidelines® NCCN-Thyroid Carcinoma [Version 3.2020]
- 30. NCCN Guidelines® NCCN-Esophageal and Esophagogastric Junction Cancers [Version 1.2021]

Date: 03 Jun 2021

References (continued)

- 31. NCCN Guidelines® NCCN-Ovarian Cancer [Version 1.2021]
- 32. NCCN Guidelines® NCCN-Breast Cancer [Version 3.2021]
- 33. https://www.focr.org/tmb
- 34. http://www.iqnpath.org/category/tmb
- 35. Stenzinger et al. Tumor mutational burden standardization initiatives: Recommendations for consistent tumor mutational burden assessment in clinical samples to guide immunotherapy treatment decisions. Genes Chromosomes Cancer. 2019 Aug;58(8):578-588. PMID: 30664300
- 36. Merino et al. Establishing guidelines to harmonize tumor mutational burden (TMB): in silico assessment of variation in TMB quantification across diagnostic platforms: phase I of the Friends of Cancer Research TMB Harmonization Project. J Immunother Cancer. 2020 Mar;8(1). PMID: 32217756