

 Ex-factory (HTVA)

 OPDIVO
 40 mg
 €509,90

 OPDIVO
 100 mg
 €1.274,75

 OPDIVO
 240 mg
 €3.059,65

Braine-l'Alleud, 1 February 2022

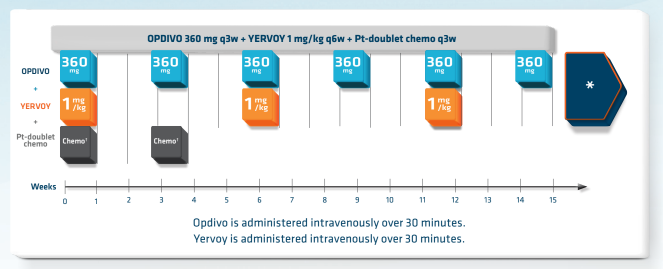
Dear Physician,

# 2022 brings us new hope for your patients.

**Bristol-Myers Squibb** is pleased to announce that both **OPDIVO**<sup>®</sup> (nivolumab) and **YERVOY**<sup>®</sup> (ipilimumab) **will be reimbursed as of 1 February** in combination with ipilimumab and 2 cycles of platinum-based chemotherapy is indicated for the first-line treatment of metastatic non-small cell lung cancer in adults whose tumours have no sensitising EGFR mutation or ALK translocation.

The recommended dose in **360 mg OPDIVO**<sup>®</sup> administered intravenously over **30 minutes** every 3 weeks in combination with **1 mg/kg YERVOY**<sup>®</sup> administered intravenously over **30 minutes** every 6 weeks, and 2 cycles of platinum-based chemotherapy administered every 3 weeks.

After completion of 2 cycles of chemotherapy, treatment is continued with **360 mg OPDIVO**<sup>®</sup> administered intravenously **every 3 weeks** in combination with **1 mg/kg YERVOY**<sup>®</sup> **every 6 weeks**.



\* Treatment is recommended until disease progression, unacceptable toxicity or up to 24 months without disease progression.

When used in combination with **YERVOY®** and/or chemotherapy, **OPDIVO®** should be given first, followed by **YERVOY®** and then chemotherapy on the same day.

Kind regards,

Gert Heymans Business Unit Director Immuno-Oncology

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Tom Van Lee Disease Area Head Immuno-Oncology

# **REIMBURSED INDICATIONS**

MONOTHERAPY		OPDIVO. (nivolumab)	Fixed dose 240 mg every 2 weeks	Fixed dose 480 mg every 4 weeks	
Melanoma <sup>a</sup>		Adjuvant	✓ 30 minutes infusion	✓ 60 minutes infusion	
Melanoma <sup>α</sup>		1L+	✓ 30 minutes infusion	✓ 60 minutes infusion	
$\begin{array}{c} \text{Renal Cell} \\ \text{Carcinoma}^{\beta} \end{array} \qquad $		2L+	✓ 30 minutes infusion	✓ 60 minutes infusion	
Non Small Cell Lung Cancer <sup>v</sup>		2L+	✓ 30 minutes infusion		
Classical Hodgkin Lymphoma <sup>8</sup>		After ASCT and BV#	✓ 30 minutes infusion		
Squamous Cell Cancer of the Head and Neck <sup>e</sup>		After failure of previous platinum-containing therapy	✓ 30 minutes infusion		
Urothelial Carcinoma <sup>ζ</sup> *		After failure of previous platinum-containing therapy	✓ 30 minutes infusion		
Oesophageal squamous cell carcinoma	A	fter fluoropyrimidine and platinum containing therapy	✓ 30 minutes infusion		
Oesophageal or gastro-oesophageal junction cancer		Adjuvant	✓ 30 minutes infusion Week 0 to 16	✓ 30 minutes infusion Week 0 to 16	
	at of troats			✓ 30 minutes infusion Week 16 to 52	
* Only for the extension of the reimbursement	OPD	OPDIVO     + YERVOY     COMBINATION PHASE     OPDIVO     MONOTHERAPY PHASE			
COMBINATION		Weight-based dosis	Fixed dose 240 mg every 2 weeks	Fixed dose 480 mg every 4 weeks	
Melanoma <sup>a</sup>	1L+	4 infusions every 3 weeks: • OPDIVO 1mg/kg 30 min infusion time • YERVOY 3mg/kg 90 min infusion time	✓ 30 minutes infusion (start after 3 weeks)	✓ 60 minutes infusion (start after 6 weeks)	
Renal Cell Carcinoma <sup>β</sup>	1L	4 infusions every 3 weeks: • OPDIVO 3mg/kg 30 min infusion time • YERVOY 1mg/kg 30 min infusion time	✓ 30 minutes infusion (start after 3 weeks)	✓ 60 minutes infusion (start after 6 weeks)	
COMBINATION COMBINATION COMBINATION PHASE					
		Week 0 to 3	Week 6	5 to 104	
Non Small Cell Lung Cancer**	1L	<ul> <li>2 cycles of platinum-based chemotherapy administered every</li> <li>3 weeks</li> <li>• OPDIVO 360 mg 30 min infusion time every 3 weeks</li> <li>• YERVOY 1 mg/kg 30 min infusion time every 6 weeks</li> </ul>	<ul> <li>OPDIVO 360 mg 30 min infusion time every 3 weeks</li> <li>YERVOY 1 mg/kg 30 min infusion time every 6 weeks</li> </ul>		
Mesothelioma***	1L	<ul> <li>• OPDIVO 360 mg 30 min infusion time every 3 weeks</li> <li>• YERVOY 1 mg/kg 30 min infusion time every 6 weeks</li> </ul>			
* Yervoy® in combination with Opdivo® in the treatment of MPM in 1 <sup>st</sup> line has not been reimbursed yet					
COMBINATION		nivolumab) + CHEMOTHERAPY	Fixed dose 240 mg every 2 weeks	Fixed dose 360 mg every 3 weeks	
Gastric, GEJ or esophageal adenocarcinoma	1L	In combination with fluoropyrimidine and platinum containing therapy	30 minutes infusion + fluoropyrimidine & platinum-based CT	30 minutes infusion + fluoropyrimidine & platinum-based CT	

\*\* Treatment is recommended until disease progression, unacceptable toxicity, or up to 24 months in patients without disease progression. Treatment with OPDIVO®, either as a monotherapy or in combination with YERVOY®, should be continued as long as clinical benefit is observed or until treatment is no longer tolerated by the patient. For adjuvant therapy, the maximum treatment duration with OPDIVO® is 12 months.



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J product is provide in program is product and program is program is product and product and product and product and product an 

dWMR or NS1H ERC, the incidence of nephrits or rend dysfunction was 8,6% (57/666). Grade 2, Grade 3, and Grade 4 cases were reported in 3.8% (52/666). 0, 0, 6% (4/666), of prients, respectively. Median time to neset was 2.1 months (range: 0.034.8), Resolution occurred in 45 patients (78.9%) with a median time to resolution of 10.0 weeks (range: 0.1-106.0°). In prients teated with nivolumob 3 rang/kg in combination with plintmumph Img/kg in XPM, the incidence of repairly (5/666), 0, 0, 6% (4/666), of prients, respectively. Median time to neset was 2.1 months (range: 0.034.8), Resolution occurred in 14 patients (78.9%) with a median time to resolution of 1.0 weeks (range: 0.1-24.4). In prients teated with nivolumob 3 rang/kg in combination with cheronherapy in gratic, EEL or oecosphagel devocarcinom, the incidence of nephritis reand dysfunction was 3.8% (5/7.666). Grade 2, Grade 3, and Grade 4 cases were reported in 2.8% (5/7.666). Grade 3, grade Grade 5 cases were reported in 2.8% (5/7.666). Grade 2, Grade 3, and Grade 4 cases were reported in 2.8% (5/7.666). Grade 2, Grade 3, and Grade 4 cases were reported in 2.8% (5/7.666). Grade 2, Grade 3, and Grade 4 cases were reported in 2.8% (5/7.666). Grade 2, Grade 3, and Grade 4 cases were reported in 2.8% (5/7.666). Grade 2, Grade 3, and Grade 4 cases were reported in 2.8% (5/7.666). Grade 2, Grade 3, and Grade 4 cases were reported in 2.8% (5/7.666). Grade 2, Grade 3, and Grade 4 cases were reported in 2.8% (3/7.8%). In prients teated with nivolumab 320 mg in combination with durammemme mediant free of second acades, including the order of the order as a case as were reported in 3.8% (2/7.358). In the order of the order as a case as were reported in 3.8% (2/7.320). Grade 2, Grade 3, and Grade 4 cases were reported in 3.8% (2/7.320). The indicates are also in a discuster as a case as were torbust 2, in the cases were reported in 0.28 (1) 2007, 2.24 (1) 2 combindion with ipilimumda 1 mg/kg every 6 weeks and chemotherapy in NSCL, the incidence of thyviaid isoardes wee 24% (86/558). Grade 2 and Grade 3 thread isoardes wee reported in 12.3% (44/358) and 0.3% (1/358) of patients, respectively. Hypophysitis occurred in 1.4% (5/558) of patients, Grade 2 and Grade 3 coses were reported in 0.4% (2/358) of patients, respectively, Made and minimum and 1 mg/kg (3/358) of patients, Grade 2 and Grade 3 coses were reported in 0.4% (2/358) of patients, respectively, Made and minimum and 1 mg/kg (3/358) of patients, Grade 2 and Grade 3 coses were reported in 0.4% (2/358) of patients, Grade 2 and Grade 3 coses were reported in 0.4% (2/358) of patients, respectively, Madian time to orset of these endocrinopathies was 12.4 and 3% (2/300) of patients, respectively, Madian time to orset of these endocrinopathies was 12.3 weeks. Competitively, Maghyphysic is courred in 0.4% (2/3200) of patients, respectively, Madian time to orset of these endocrinopathies was 12.3 weeks. Competitively, Maghyphysic is courred in 0.4% (2/3200) of patients, respectively, Madian time to orset of these endocrinopathies was 12.3 weeks. Competitively, Maghyphysic is courred in 0.4% (2/3200) of patients, respectively, Madian time to orset of 1 a service reported in 2.2% (1/220) ond 1.9% (6/320) of patients, respectively, Madian time to orset of 1 a service reported in 2.2% (1/220) ond 1.9% (6/320) of patients, respectively, Madian time to orset of 1 a service reported in 2.2% (1/220) ond 1.9% (6/320) of patients, respectively, Madian time to orset of response 3.4% (1/27/371). In patients thereted with involumed 3 mg/kg in mediance of resh was 5.5% (29/1/440) or patients, respectively, Madian time to orset was 1.0 months (range: 0.19.4% (20) and 1.1% (9/300) of patients, respectively, Madian time to orset was 1.0 months (range: 0.019.4% (20) and 1.1% (9/300) of patients, respectively, Madian time to orset was 1.0 months (range: 0.019.4% (20) and 1.2% (20 weeks: compact U-130.c weeks), kine coses of S.S. on tell some of them with thord actione there been doesrved (see schools + 2 and + 3.1. //listication) //micro mechanics weeks and tell some of them with thord actione there been doesrved (see schools + 2 and + 3.1. //listication) //micro mechanics weeks and tell some of them with molumab 1 mg/kg in combination with planmamb 1 mg/kg in planets treated with molumab 3 mg/kg in combination with planmamb 1 mg/kg in the does mg/kg in combination with planmamb 1 mg/kg in the does mg/kg in combination with planmamb 1 mg/kg in the does mg/kg in combination with planmamb 1 mg/kg in the does mg/kg in combination with planmamb 1 mg/kg in the does mg/kg in combination with planmamb 1 mg/kg in the does mg/kg in combination with planmamb 1 mg/kg in the does mg/kg in combination with planmamb 1 mg/kg in the does mg/kg in combination with planmamb 1 mg/kg in the does mg/kg in combination with planmamb 1 mg/kg in the does mg/kg in combination with planmamb 1 mg/kg in the does wee reported in 2.4% (16/666) of patients. Respectively, In patients treated with molumab 3 ang/kg in combination with planmamb 1 mg/kg in the does wee reported in 2.4% (16/666) of patients. Respectively, In patients treated with molumab 3 ang/kg in combination with planmamb 1 mg/kg in the does wee reported in 2.4% (10/480) of patients. Respectively, In patients treated with molumab 3 ang/kg in combination with charantherapy in gastin, GEI or escolution of 0.1 weeks (nonge; 0.1 ?9.1). In patients treated with molumab 3 mg/kg in excellulates weeks and a set of the does and excellulates weeks and a set of the does and excellulates weeks and a set of the does and excellulates and a set of the does and ex cell infusion, was reported in four patients (6%). A steriot-requiring fabrile syndrome, without an identified infectious cause, was reported in six patients (12%) within the first 6 weeks post-transplointation. Steriods were used in four patients on d three patients responded to steriods. Hepatic vene-occlusive disease accurred in two patients, one of whom died of GWD and multi-arguin fabrice. Nineteen of 62 patients (30.6%) died from complications of allogeneic HSCT ofter nivolumab. The 62 patients had a median follow-up from subsequent allogeneic HSCT of 38.5 months (range: 0+8 months). *Elevated liver enzymes when nivolumab is combined with cabozantinib. in RCC* In a clinical study of previously untreated patients with RC receiving to planter the construction of the constructio in 2 potents reaving CPUID 4, petites reaving calcaratini, and 2 priorits reaving calcaratini, and 2 p Medicinal product subject to restricted medical prescription 11. DATE OF REVISION OF THE TEXT 07 December 2021 Detailed information on this medicinal product is available on the website of the European Medicines Agency http://www.ema.europa.eu

### « MELANOMA

OPDIVO as monotherapy is indicated for the adjuvant treatment of adults with melanoma with involvement of lymph nodes or metastatic disease who have undergone complete resection

OPDIVO as monotherapy or in combination with ipilimumab is indicated for the treatment of advanced (unresectable or metastatic) melanoma in adults. Relative to nivolumab monotherapy, an increase in progression-free survival (PFS) and overall survival (OS) for the combination of nivolumab with ipilimumab is established only in patients with low tumour PD-L1 expression Permonizional as monotherapy is indicated for the treatment of advanced (unresectable or metastatic) melanoma in adults P RENAL CELL CARCINOMA

OPDIVO as monotherapy is indicated for the treatment of advanced renal cell carcinoma after prior therapy in adults.

### NON SMALL CELL LUNG CANCER

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OPDIVO as monotherapy is indicated for the treatment of locally advanced or metastatic non-small cell lung cancer after prior chemotherapy in adults Pembrolizumab, as monotherapy is indicated in the treatment of locally advanced or metastatic NSCLC in adults whose tumours express PD-L1 with a 21% TPS and who have received at least one prior chemotherapy. Patients with EGFR or ALK positive tumour mutations should also have received targeted therapy before receiving pembrolizumab.

## Atezoliziumab is monotherapy is indicated for the treatment of locally advanced or metastatic non-small cell lung cancer (NSCLC) in adults after prior chemotherapy. Patients with EGFR or ALK positive tumour mutations should also have received targeted therapy before receiving atezolizumab.

° CLASSICAL HODGKIN LYMPHOMA

Depuive a monotherapy is indicated for the treatment of adult patients with relapsed or refractory classical Hodgkin lymphoma after autologous stem cell transplant (ASCT) and treatment with brentuximab vedotin. Pembrolizumab as monotherapy is indicated in the treatment of adult patients with relapsed or refractory classical Hodgkin (cHL) who failed autologous stem cell transplant (ASCT) and brentuximab vedotin.

E HEAD AND NECK

OPDIVO as monotherapy is indicated for the treatment of recurrent or metastatic squamous cell cancer of the head and neck in adults progressing on or after platinum-based therapy. Keytruda as monotherapy is indicated for the treatment of recurrent or metastatic head and neck squamous cell carcinoma (HNSCC) in adults whose tumours express PD-L1 with a 2 50% TPS and progressing on or after platinum-containing chemotherapy **CUROTHELIAL CARCINOMA** 

OPONDO as monotherapy is indicated for the treatment of locally advanced unresectable or metastatic urothelial carcinoma in adults after failure of prior platinum-containing therapy Pembrolizumab as monotherapy is indicated in the treatment of adult patients with locally advanced or metastatic urothelial carcinoma in adults who have received prior platinum-based therapy. Pembrolizumab as monotherapy is indicated in the treatment of adult patients with locally advanced or metastatic urothelial carcinoma in adults who have received prior platinum-based therapy. Pembrolizumab as monotherapy is indicated in the treatment of adult patients with locally advanced or metastatic urothelial carcinoma in adults who have received prior platinum-based therapy. Pembrolizumab as monotherapy is indicated in the treatment of adult patients with locally advanced or metastatic urothelial carcinoma in adults who have received prior platinum-based therapy. Pembrolizumab as monotherapy is indicated in the treatment of adult patients with locally advanced or metastatic urothelial carcinoma in adults who have received prior platinum-based therapy. Pembrolizumab as monotherapy is indicated in the treatment of adult patients with locally advanced or metastatic urothelial carcinoma in adults who have received prior platinum-based therapy. Pembrolizumab as monotherapy is indicated in the treatment of adult patients with locally advanced or metastatic urothelial carcinoma in leigible for cisplatin-based chemotherapy and whose tumours express PD-L1 with a combined positive score (CPS) is 10. Atezolizumab as monotherapy is indicated for the treatment of adult patients with locally advanced or metastatic urothelial carcinoma

after one prior platinum-based chemotherapy, or - considered ineligible for cisplatin and whose tumours show PD-L1 expression ≥ 5%