

Immunotherapy for the Treatment of Lung Cancer

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Society for Immunotherapy of Cancer





- Consulting Fees:
 - Genentech/Roche, Novartis, Celgene, BMS, AstraZeneca, Pfizer, Boehringer Ingelheim, Abbvie, Foundation Medicine, GlaxoSmithKline, Lilly, Merck, Moderna Therapeutics, Nektar, Takeda, Amgen, TRM Oncology, Precision Oncology, Evelo Therapeutics, Illumina, PharmaMar; Contracted Research: Genentech/Roche, Novartis, Celgene, BMS, Lilly, AstraZeneca, Pfizer, Boehringer Ingelheim, UT Southwestern Medical Center - Simmons Cancer Center, Merck, Abbvie, GlaxoSmithKline, G1 Therapeutics, Neon Therapeutics, Takeda, Foundation Medicine, Nektar, Celldex, Clovis Oncology, Daiichi Sankyo, EMD Serono, Acerta Pharma, Oncogenex, Astellas Pharma, GRAIL, Transgene, Aeglea Biotherapeutics, Tesaro, Ipsen, ARMO BioSciences, Amgen, Millennium
- I will be discussing non-FDA approved indications during my presentation.





- 80-85% non-small cell lung cancer (NSCLC)
- 10-15% small cell lung cancer (SCLC)
- NSCLC has relatively long and extensive history of immunotherapy use

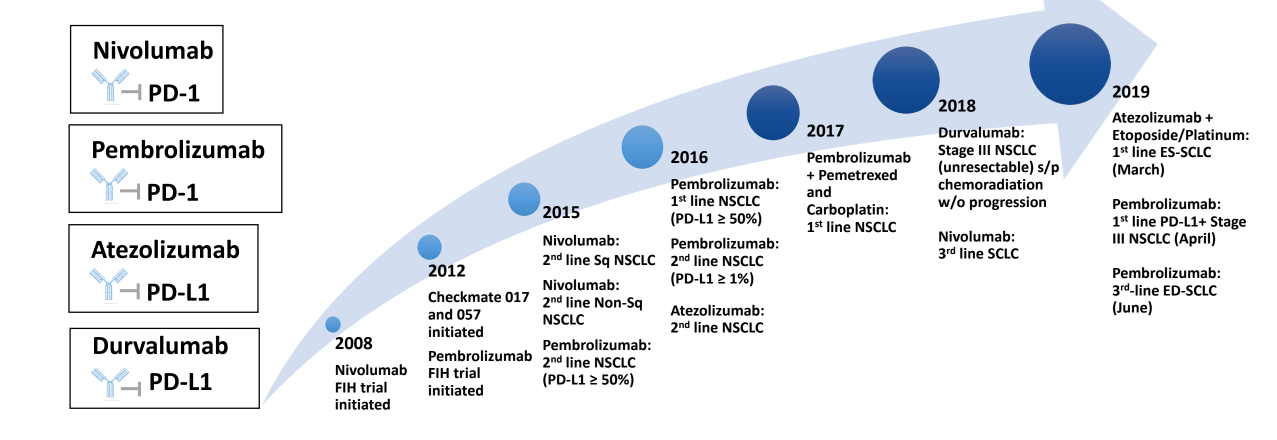
	Male			
	Lung & bronchus	76,650	24%	
	Prostate	31,620	10%	
s	Colon & rectum	27,640	9%	
ath	Pancreas	23,800	7%	
Estimated Deaths	Liver & intrahepatic bile duct	21,600	7%	
	Leukemia	13,150	4%	
	Esophagus	13,020	4%	
<u>a</u> .	Urinary bladder	12,870	4%	
ES1	Non-Hodgkin lymphoma	11,510	4%	
	Brain & other nervous system	9,910	3%	
	All sites	321,670		

remate		
Lung & bronchus	66,020	23%
Breast	41,760	15%
Colon & rectum	23,380	8%
Pancreas	21,950	8%
Ovary	13,980	5%
Uterine corpus	12,160	4%
Liver & intrahepatic bile duct	10,180	4%
Leukemia	9,690	3%
Non-Hodgkin lymphoma	8,460	3%
Brain & other nervous system	7,850	3%
All sites	285,210	

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FDA-approved checkpoint inhibitors in lung cancer





Approved checkpoint inhibitors in NSCLC

Drug	Approved	Indication	Dose
	2015	Metastatic Squamous NSCLC with progression after chemotherapy (2 nd line)	240 mg Q2W or 480 mg
Nivolumab	2015	Metastatic Non-Squamous NSCLC with progression after chemotherapy (2 nd line)	Q4W





Approved checkpoint inhibitors in NSCLC

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	Drug	Approved	Indication	Dose
		2015	Metastatic NSCLC with progression after chemotherapy and PD-L1 ≥ 50%	
		2016	Metastatic NSCLC with progression after chemotherapy and PD-L1 ≥ 1%	
	Pembrolizumab	2016	1^{st} line metastatic NSCLC with PD-L1 TPS \ge 50%	
		2019	1 st line stage III NSCLC (not candidate for resection or definitive chemoradiation) and Metastatic NSCLC, with PD-L1 TPS ≥ 1% and no EGFR/ALK mutations	200 mg Q3W
	Pembrolizumab + pemetrexed & carboplatin	2017	1 st line metastatic Non-Squamous NSCLC	
	Pembrolizumab + pemetrexed + platinum	2018	1 st line metastatic Non-Squamous NSCLC with no EGFR/ALK mutations	
© 2	Pembrolizumab + carboplatin + paclitaxel/nab-paclitaxel	2018	1 st line metastatic Squamous NSCLC	



Approved checkpoint inhibitors in NSCLC

Drug	Approved	Indication	Dose
Atezolizumab	2016	Metastatic NSCLC with progression after Pt-chemotherapy and targeted therapy if EGFR/ALK mutation-positive	840 mg Q2W, 1200 mg Q3W, or 1680 mg Q4W
Atezolizumab + bevacizumab + paclitaxel + carboplatin	2018	1 st line metastatic non-squamous NSCLC with no EGFR/ALK mutations	For 4-6 cycles: atezolizumab 1200 mg Q3W + chemotherapy + bevacizumab Maintenance: 840 mg Q2W, 1200 mg Q3W, or 1680 mg Q4W
Durvalumab	2018	Stage III NSCLC, ineligible for surgery and without progression after chemoradiation	10 mg/kg Q2W





Treatment Naïve Regimens: Competing Strategies in NSCLC

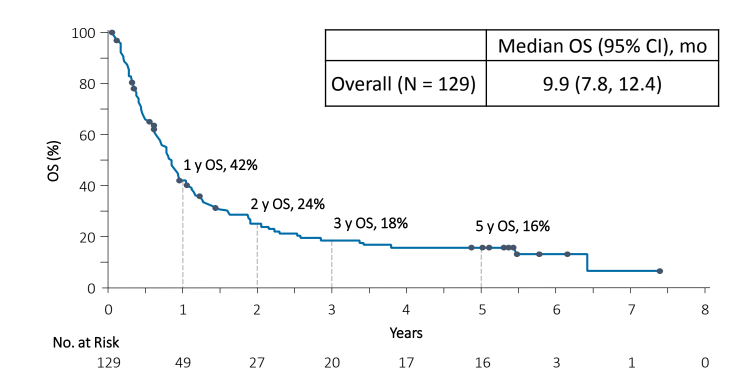
- **KEYNOTE 024** Pembrolizumab vs. Chemotherapy in PD-L1 ≥ 50%
- **KEYNOTE 042** Pembrolizumab vs. Chemotherapy in PD-L1 ≥ 1%
- KEYNOTE 189 Pembrolizumab + Chemotherapy vs. Chemotherapy alone in advanced non-squamous NSCLC
- IMPOWER 150 Atezolizumab + Chemotherapy (Bev) vs. Chemotherapy (Bev) in advanced non-squamous NSCLC
- KEYNOTE 407 Pembrolizumab + Chemotherapy vs. Chemotherapy in advanced squamous cell lung cancer
- CHECKMATE 227 Ipilimumab + Nivolumab vs. Chemotherapy in advanced NSCLC with high TMB





CA209-003: Nivolumab in Heavily-pretreated Advanced NSCLC (NCT00730639) Phase 1, 5-Year Update 5-Year Survival

- First report of long-term survival rate in patients with metastatic NSCLC treated with an immune checkpoint inhibitor
- According to the National Cancer Institute's SEER data, 5-year survival rate for patients with advanced NSCLC is 4.9%



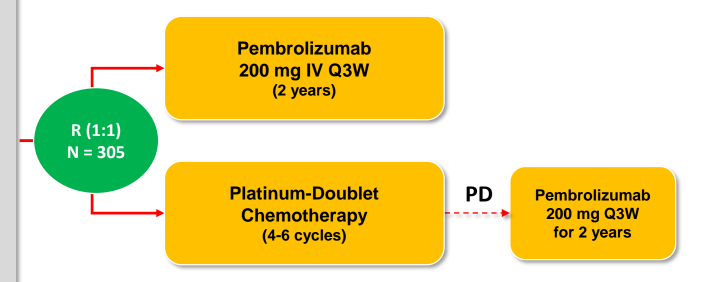




KEYNOTE-024: Pembrolizumab vs. Chemotherapy for PD-L1 Positive (≥ 50%) NSCLC Study Design (NCT021427389)



- Untreated stage IV NSCLC
- PD-L1 TPS ≥50%
- ECOG PS 0-1
- No activating *EGFR* mutation or *ALK* translocation
- No untreated brain metastases
- No active autoimmune disease requiring systemic therapy



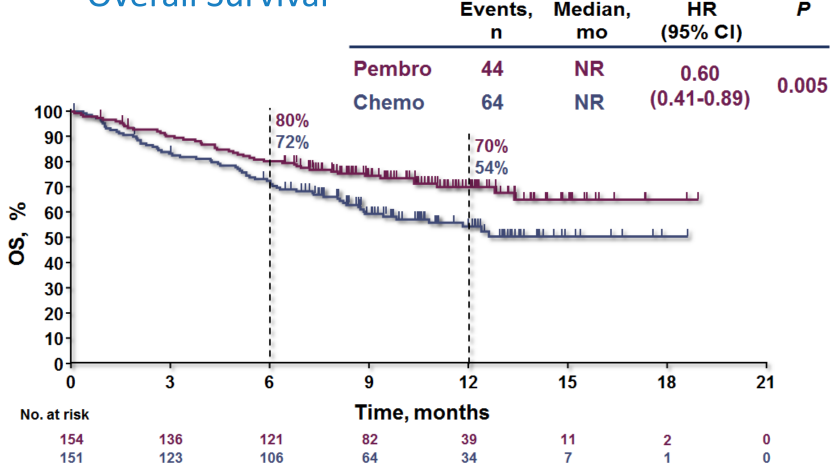




KEYNOTE-024: Pembrolizumab vs. Chemotherapy for PD-L1 ≥ 50% NSCLC Overall Survival

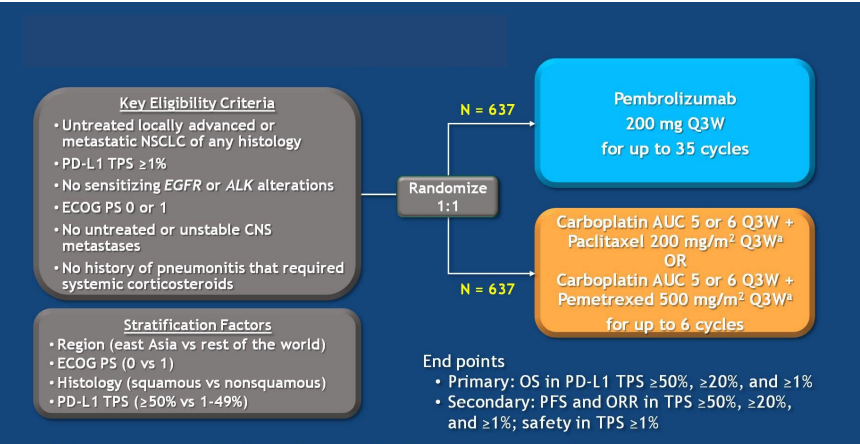
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KEYNOTE-042: Pembrolizumab vs. Chemotherapy for PD-L1 ≥ <u>1%</u> NSCLC

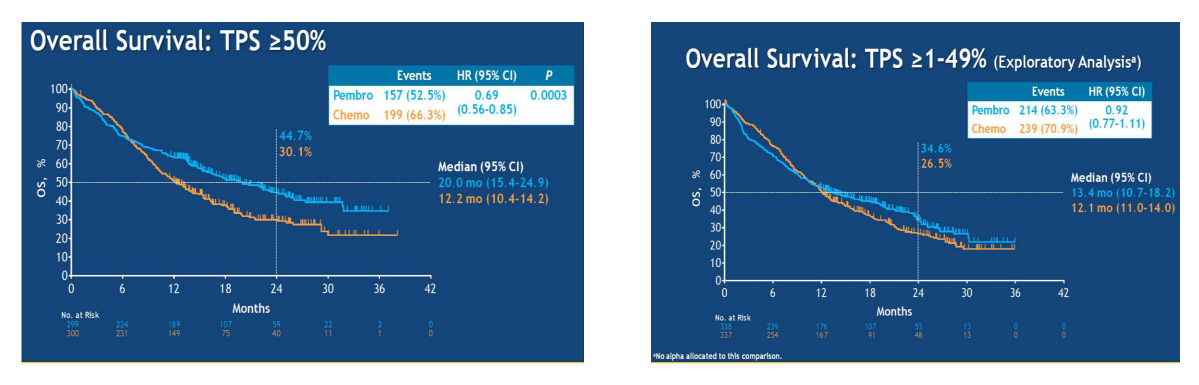


^aPemetrexed maintenance therapy was optional but strongly encouraged for patients with nonsquamous histology.





KEYNOTE-042: Pembrolizumab vs. Chemotherapy for PD-L1 ≥ 1% NSCLC Overall Survival



Survival benefit seemed to be driven by the TPS ≥ 50% subset with little benefit witnessed in the subset TPS = 1 - 49%





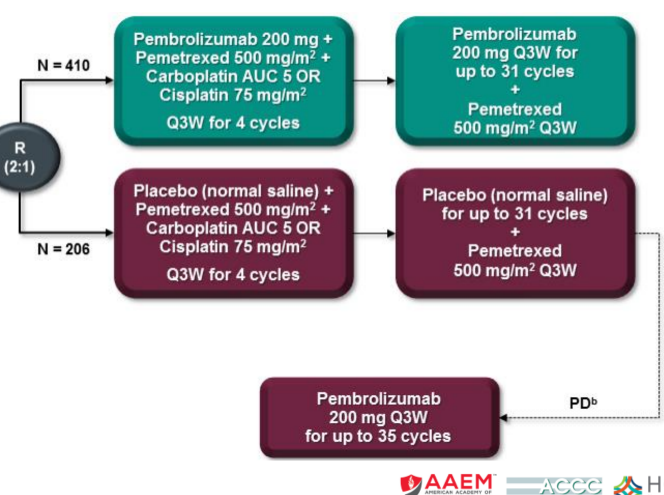
KEYNOTE-189: Pembrolizumab/Platinum /Pemetrexed vs Chemotherapy Alone for Advanced Non-Squamous NSCLC

Key Eligibility Criteria

- Untreated stage IV nonsquamous NSCLC
- No sensitizing EGFR or ALK alteration
- ECOG PS0 or 1
- Provision of a sample for PD-L1 assessment
- No symptomatic brain metastases
- No pneumonitis requiring systemic steroids

Stratification Factors

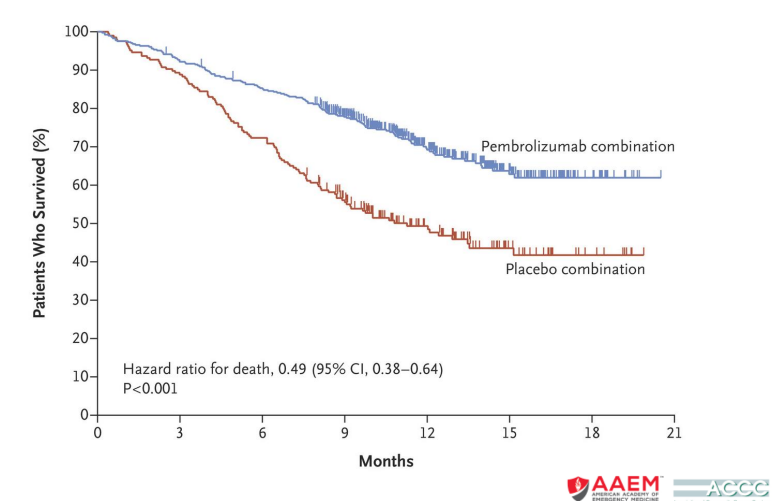
- PD-L1 expression (TPS^a <1% vs ≥1%)
- Platinum (cisplatin vs carboplatin)
- Smoking history (never vs former/current)



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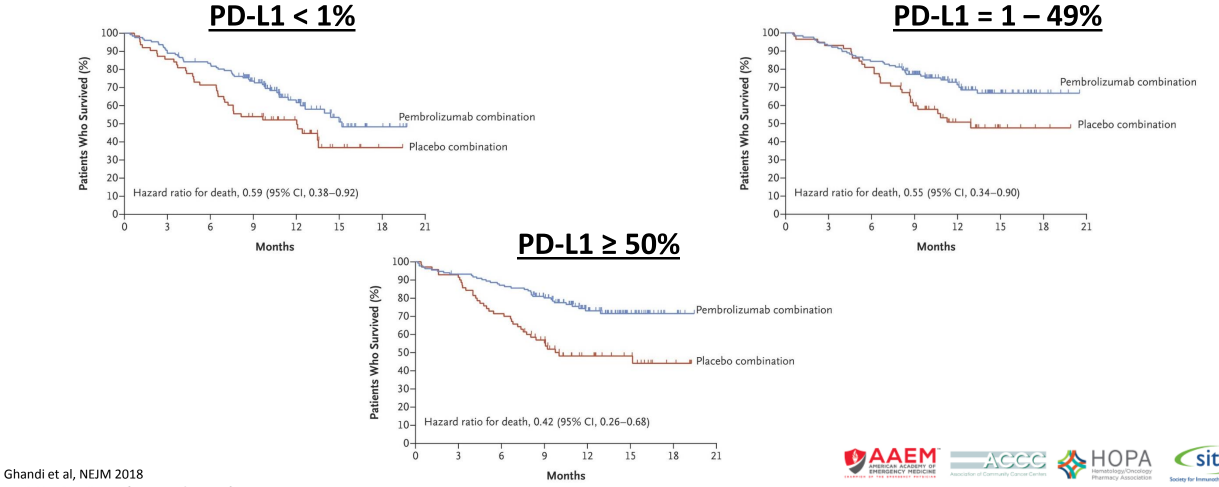


KEYNOTE-189: Pembrolizumab/Platinum /Pemetrexed vs Chemotherapy Alone for Advanced Non-Squamous NSCLC





KEYNOTE-189: Pembrolizumab/Platinum /Pemetrexed vs Chemotherapy Alone for Advanced Non-Squamous NSCLC



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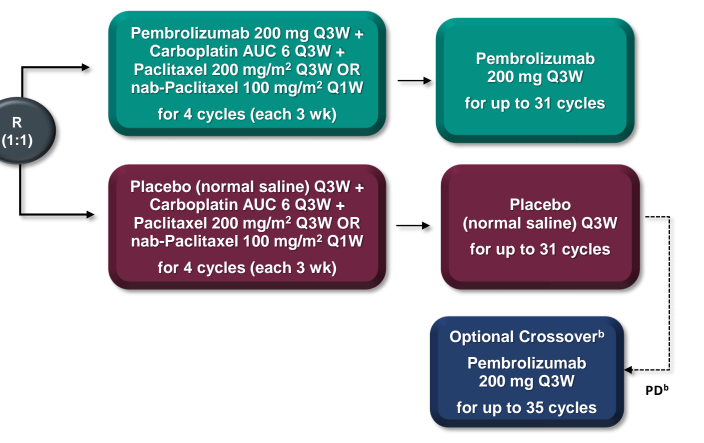
KEYNOTE-407: Pembrolizumab/Chemotherapy vs Chemotherapy Alone for Advanced Squamous-Cell NSCLC

Key Eligibility Criteria

- Untreated stage IV NSCLC with squamous histology
- ECOG PS 0 or 1
- Provision of a sample for PD-L1 assessment
- No symptomatic brain metastases
- No pneumonitis requiring systemic steroids

Stratification Factors

- PD-L1 expression (TPS^a <1% vs ≥1%)
- Choice of taxane (paclitaxel vs nab-paclitaxel)
- Geographic region (east Asia vs rest of world)

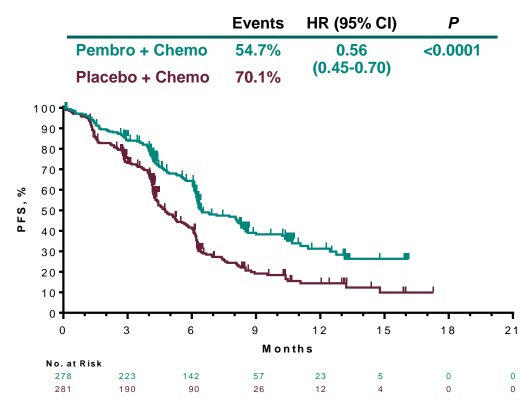


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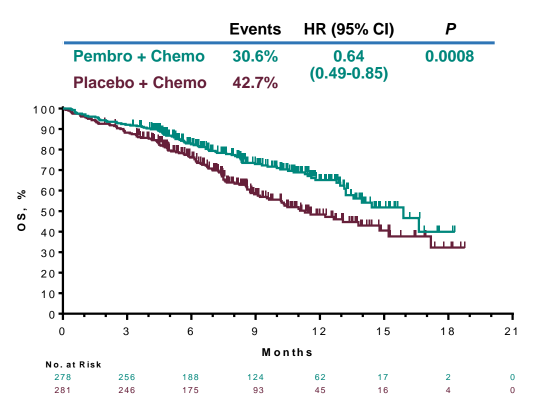


KEYNOTE-407: Pembrolizumab/Chemotherapy vs Chemotherapy Alone for Advanced Squamous-Cell NSCLC

PFS (RECISTv1.1, BICR)



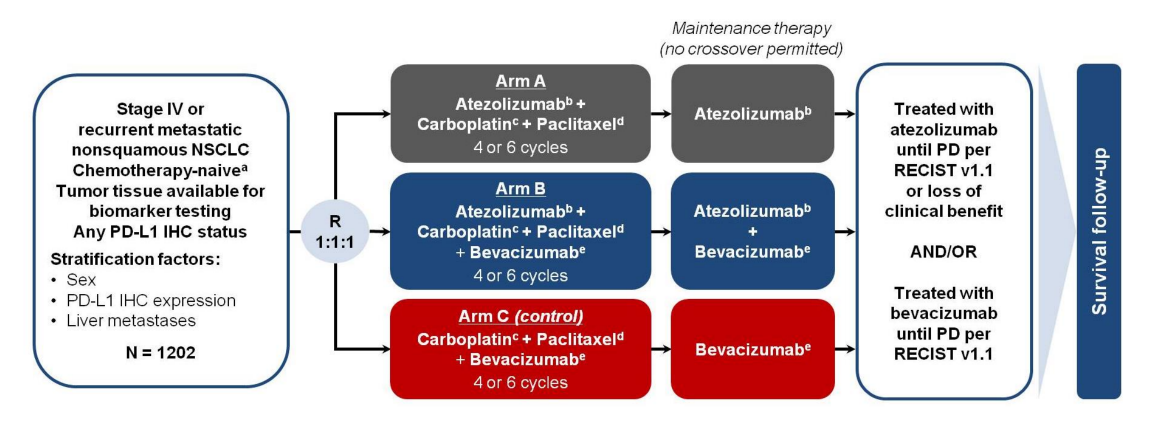
Overall Survival







IMPOWER 150: Atezolizumab/Carboplatin/ Paclitaxel/Bevacizumab vs Carboplatin/Paclitaxel/ Bevacizumab in Advanced Non-Squamous NSCLC

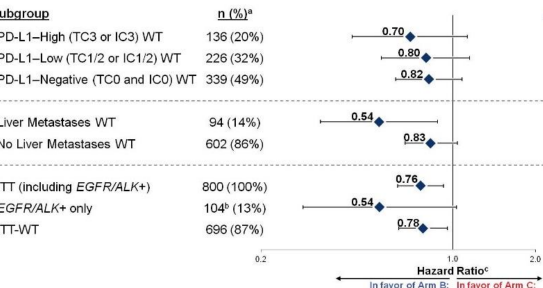


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IMPOWER 150: Atezolizumab/Carboplatin/ Paclitaxel/Bevacizumab vs Carboplatin/Paclitaxel/ Bevacizumab in Advanced Non-Squamous NSCLC

Landmark OS, %	Arm B: atezo + bev + CP	Arm C: bev + CP		
12-month	67%	61%	HRª, 0.78	
18-month	53%	41%	(95% CI: 0.64, 0.96) P = 0.0164	Subgroup
24-month	43%	34%	Median follow-up: ~20 mo	PD-L1—High (TC3 or PD-L1—Low (TC1/2 o
				PD-L1–Negative (TC
and and a second				Liver Metastases WT
and the second s	and			No Liver Metastases
	A DECEMBER OF THE OWNER OWNER OF THE OWNER O			ITT (including EGFR
		Statement and	200	EGFR/ALK+ only
		Row Street Stree	and the second s	ITT-WT
	1		Here with	
	edian, 14.7 mo % CI: 13.3, 16.9)		ian, 19.2 mo Cl: 17.0, 23.8)	



atezo + bev + CP bev + CP

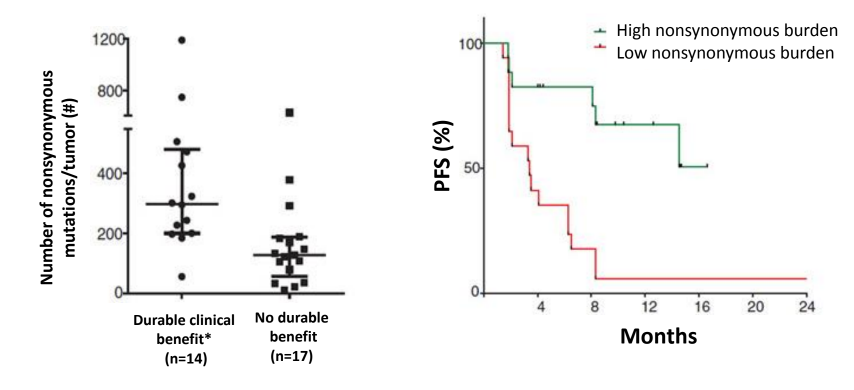


Socinski et al, NEJM 2018 © 2019–2020 Society for Immunotherapy of Cancer



Tumor Mutational Burden (TMB) may Determine Sensitivity to PD-1 Blockade in NSCLC

In two independent cohorts, higher nonsynonymous tumor mutational burden (TMB) was associated with improved objective response, durable clinical benefit, and PFS.

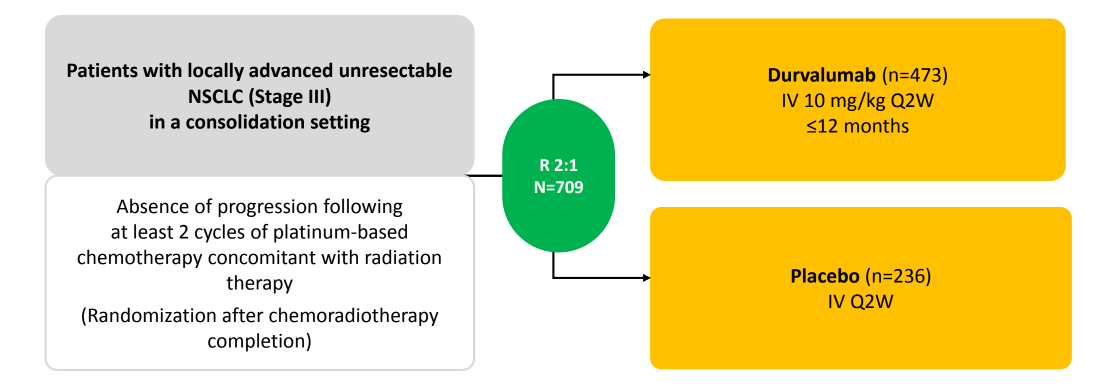


*Partial or stable response lasting > 6 mo





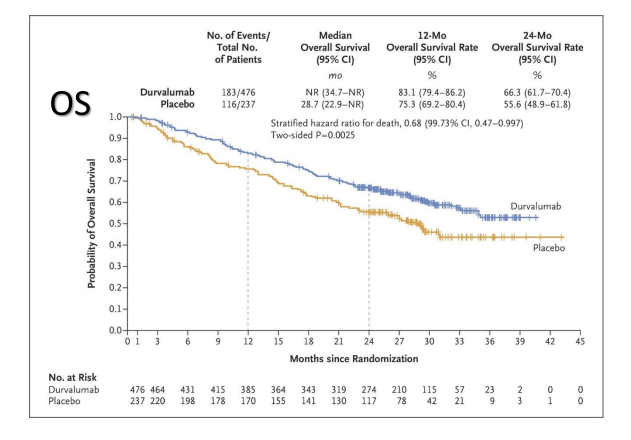
PACIFIC (NCT02125461): Durvalumab after chemoradiotherapy in Stage III NSCLC

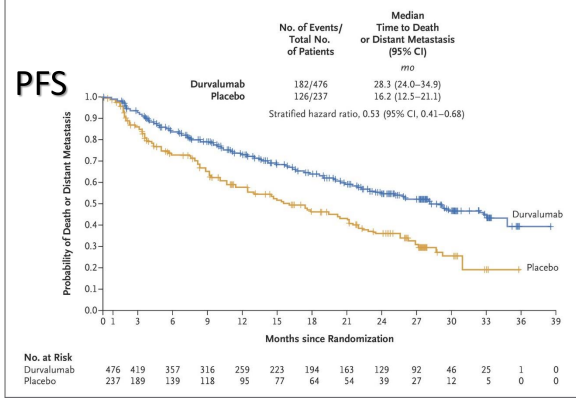






PACIFIC (NCT02125461): Durvalumab after chemoradiotherapy in Stage III NSCLC



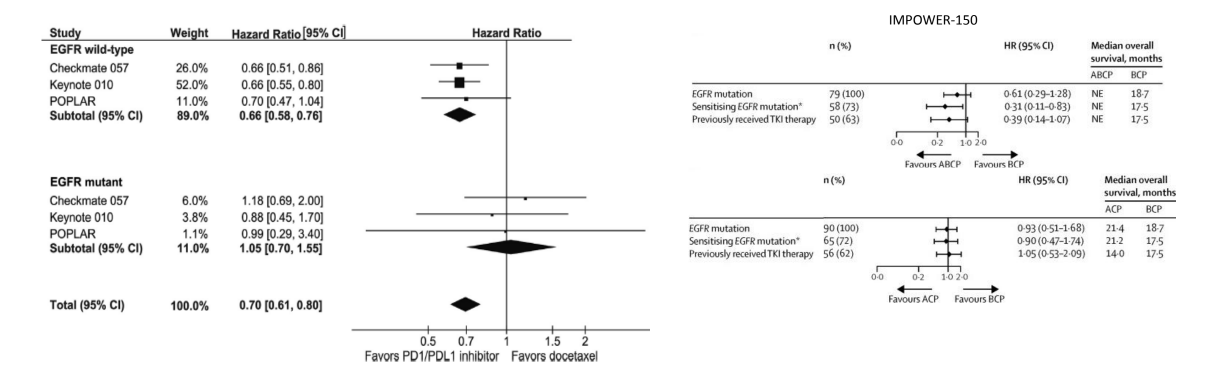






Checkpoint Inhibitors in Metastatic EGFR-Mutated NSCLC

Meta-Analysis: CM-057, KN-010, POPLAR; IMPOWER-150



CK Lee et al., JTO 2016 M Reck et al., Lancet Resp Med 2019 © 2019–2020 Society for Immunotherapy of Cancer





PD-1/PD-L1 Inhibitors Increase Overall Survival in 2L Advanced NSCLC

CHECKMATE 017 (nivolumab)	Nivolumab (N–135) Docetaxel (N–137)		 1-Yr Overall \$ % of patients (42 (34-5 24 (17-3)) 	95% CI) 0)	No. of Deaths 86 113
CHECKMATE 057 (nivolumab)	mOS, mo	Nivolumab (n = 292) 12.2	Docetaxel (n = 290) 9.4		
	HR = 0.73 (S	96% CI: 0.59, 0.89); P Median (95% CI), mo	= 0.0015 HR* (95% CI)	Р	_
(NOTE 010 (TPS ≥ 1%) (pembrolizumab)	Pembro 2 mg/kg Pembro 10 mg/kg	14.9 (10.4-NR) 17.3 (11.8-NR)	0.54 (0.38-0.77) 0.50 (0.36-0.70)	0.0002 <0.0001	_
(periorenzanab)	Docetaxel	8.2 (6.4-10.7)	-		_
OAK	HR, 0.73 ^a (95% Cl, 0.62, P = 0.0003	0.87)			

(atezolizumab)

KEYNO

Minimum follow up = 19 months



Borghaei, NEJM 2015 Herbst Lancet 2016 Rittmeyer Lancet 2017 © 2019–2020 Society for Immunotherapy of Cancer

Brahmer NEJM 2015



Small cell lung cancer

- 10-15% of lung cancers
- Almost exclusively former/current smokers
- Median survival 1-2 years after diagnosis
- Until recently, only one FDA-approved 2nd line option: topotecan DOR: 3.3 months
- Recent approvals of immunotherapies mark the first progress in decades





Approved checkpoint inhibitors in **SCLC**

Drug	Approved	Indication	Dose
Nivolumab	2018	Metastatic small cell lung cancer with progression on Pt-chemotherapy and one other therapy (3 rd line)	240 mg Q2W
Atezolizumab + carboplatin + etoposide	2019	1 st line extensive stage SCLC	For 4 cycles: atezolizumab 1200 mg + carboplatin + etoposide Q3W Maintenance: 840 mg Q2W, 1200 mg Q3W, or 1680 mg Q4W
Pembrolizumab	2019	Metastatic small cell lung cancer with progression on Pt-chemotherapy and one other therapy (3 rd line)	200 mg Q3W



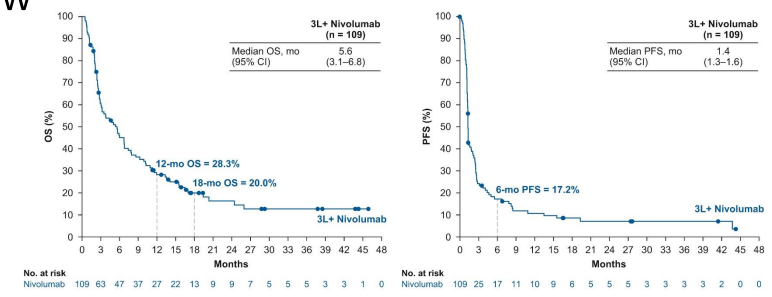






CheckMate-032: Nivolumab in 3rd line SCLC

- Nivolumab in SCLC with progression on platinum chemotherapy and another therapy
- Nivolumab 3 mg/kg Q2W
- @28.3 months:
 - ORR: 11.9%
 - mDOR: 17.9 months



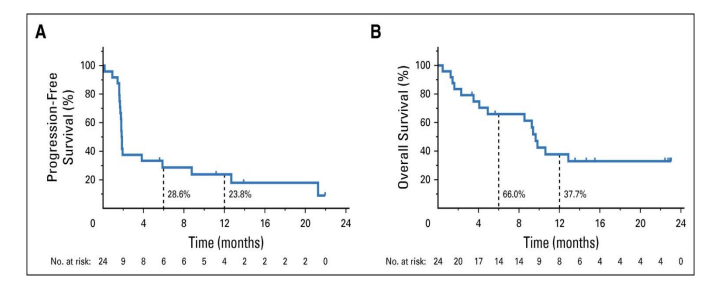




Pembrolizumab in 3rd-line SCLC

- KEYNOTE-028: PD-L1+ only (Cohort C1)
- KEYNOTE-158: PD-L1 +/-(Cohort G)
- Combined analysis:
- ORR: 19.3%
 - 2 CR, 14 PR
 - 14/16 responders were PD-L1+
 - 9/16 responders had response ≥18 mo.
- mOS: 7.7 months

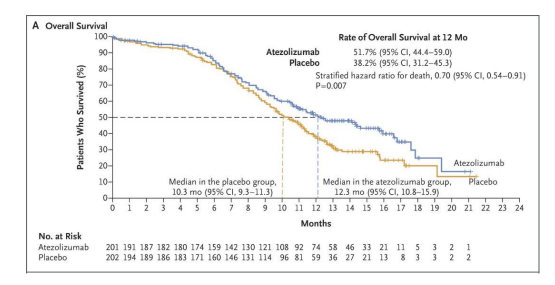






IMpower133: Atezolizumab + chemo in 1st-line SCLC

- Induction phase: four 21-day cycles of carboplatin and etoposide + atezolizumab (1200 mg once per cycle) or placebo
- Maintenance phase: either atezolizumab or placebo
- @13.9 mo:
 - mOS = 12.3 vs 10.3 mo
 - mPFS = 5.2 vs 4.3 mo









- NSCLC has been a proving ground for checkpoint inhibitors
- Moving from 2nd/3rd line options to the front line
- Clear-cut biomarkers still lacking









Brahmer et al. Journal for ImmunoTherapy of Cancer (2018) 6:75 Journal for ImmunoTherapy https://doi.org/10.1186/s40425-018-0382-2 of Cancer **POSITION ARTICLE AND GUIDELINES Open Access** CrossMark The Society for Immunotherapy of Cancer consensus statement on immunotherapy for the treatment of non-small cell lung cancer (NSCLC) Julie R. Brahmer¹, Ramaswamy Govindan², Robert A. Anders³, Scott J. Antonia⁴, Sarah Sagorsky⁵, Marianne J. Davies⁶, Steven M. Dubinett⁷, Andrea Ferris⁸, Leena Gandhi⁹, Edward B. Garon¹⁰, Matthew D. Hellmann¹¹, Fred R. Hirsch¹², Shakuntala Malik¹³, Joel W. Neal¹⁴, Vassiliki A. Papadimitrakopoulou¹⁵, David L. Rimm¹⁶, Lawrence H. Schwartz¹⁷, Boris Sepesi¹⁸, Beow Yong Yeap¹⁹, Naiyer A. Rizvi²⁰ and Roy S. Herbst^{21*}





Case Studies





Case Study 1

- Ms. AB is a 72yo retired teacher who presented in 2013 with Stage IIIB NS-NSCLC
 - No PMH otherwise
 - active smoker
 - KRAS and FGFR4 mutations
 - received chemoRT (and consolidation chemotherapy) ending 4/2013
 - she developed progressive disease in her right inguinal region 12/2013
 - started avelumab (on protocol; an anti-PD-L1 inhibitor)
 - achieved a CR
 - remains on therapy w/o toxicity





Case Study 2

- Mr. JL presented w/LS-SCLC and was treated with platinum/etoposide + RT ending 9/2013
- Relapsed disease confirmed in the liver in 2015, and went on study with nivolumab/ipilimumab 1/2015
- He had to stop all treatment due to a refractory rash/pruritus 7/2016
- He has no evidence of disease and I see twice a year

