

Approach to NSCLC Without Targetable Mutations

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First Line Lung Cancer Therapy with no actionable genes

NSQCC:

Carboplatin/Pemetrexed/Pembrolizumab [Keynote 189]

Carboplatin/Paclitaxel/Bevacizumab/Atezolizumab [IMPOWER 150]

SQCC:

Carboplatin/Paclitaxel or nab-paclitaxel/Pembrolizumab [Keynote 407]

NSQCC and SQCC:

Cemiplimab/Chemotherapy [Empower Lung-3]

Durvalumab +Tremelimumab/Chemotherapy [Poseidon 3]

IO single Agent (NSQCC OR SQCC)

Pembrolizumab [Keynote 024 and 042]

Atezolizumab [IMPOWER 110]

• Cemiplimab [Empower Lung-1]

Immunotherapy combinations:

Pilimumab and Nivolumab [Checkmate 227]

• Ipilimumab and Nivolumab plus 2 cycles of chemotherapy [Checkmate 9LA]





Follow-up

EMPOWER-Lung 3 (Part 2) Study Design (NCT03409614)

Background: Cemiplimab (a high-affinity, fully human anti–PD-1) is approved as first-line monotherapy for advanced NSCLC with PD-L1 ≥50% (EMPOWER-Lung 1 Study¹)

Key eligibility criteria

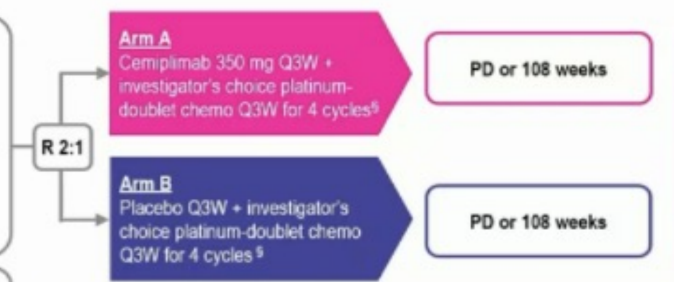
- Treatment-naive advanced NSCLC (non-squamous and squamous histology; Stage IIIb/c:, IV)
- Any PD-L1 expression
- No EGFR, ALK, or ROS1 mutations
- . ECOG PS 0 or 1
- Treated, clinically stable CNS metastases [‡]

Stratification factors

- PD-L1 expression: <1% vs 1-49% vs ≥50%
- Histology: non-squamous vs squamous

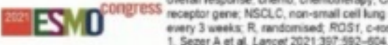
Endpoints

- Primary: OS
- Key secondary: PFS and ORR
- Additional secondary: DOR, BOR, safety, and PRO



N=466

Two interim analyses were prespecified per protocol Second interim analysis (14 June 2021) presented here



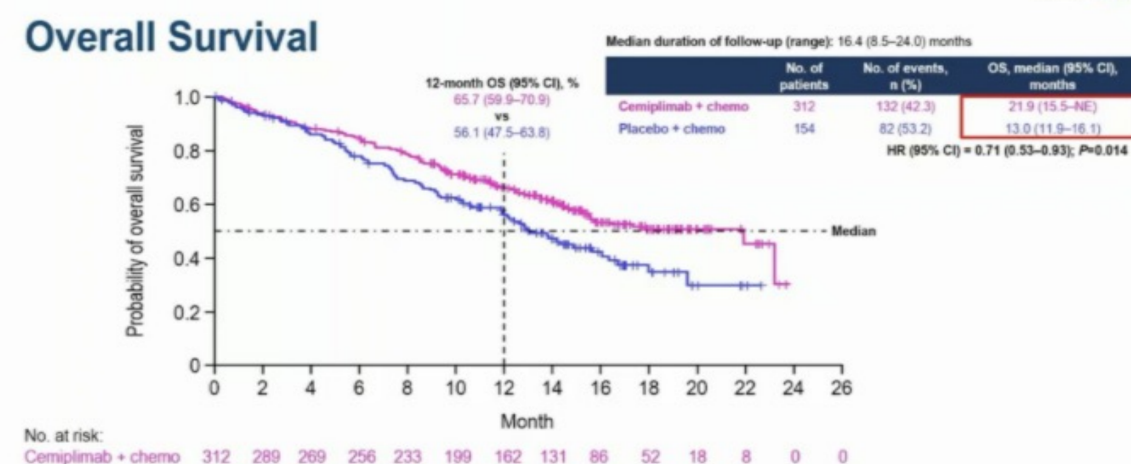
Patient not a candidate for definitive chemoradiation. ¹ Patient must have neurologically returned to baseline (except for residual signs or symptoms related to the CNS treatment). ¹For patients with non-squamous NSCLC, pernetreived is mandatory as maintenance therapy for those patients initially assigned to receive a pernetreived-containing regimen. ALK anaplastic lymphoma kinase gene; BOR, best overall response; chemo, chemotherapy; CNS, central nervous system; DOR, duration of response; ECCG PS, Eastern Cooperative Oncology Group performance status; EGFR, epidermal growth factor receptor gene; NSCLC, non-small cell lung cancer; ORR, objective response rate; OS, overall survival; PD, progressive disease; PFS, progression-free survival; PRO, patient-reported outcomes; Q3W, every 3 weeks; R, randomised; ROS1, c-los oncogene 1.

14

26









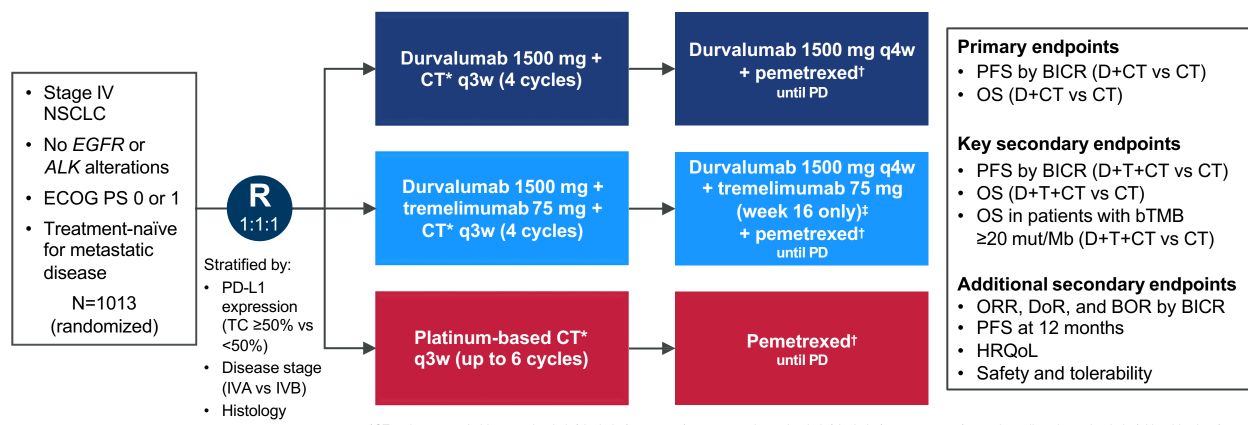
Placebo + chemo

126

112

POSEIDON Study Design

Phase 3, global, randomized, open-label, multicenter study

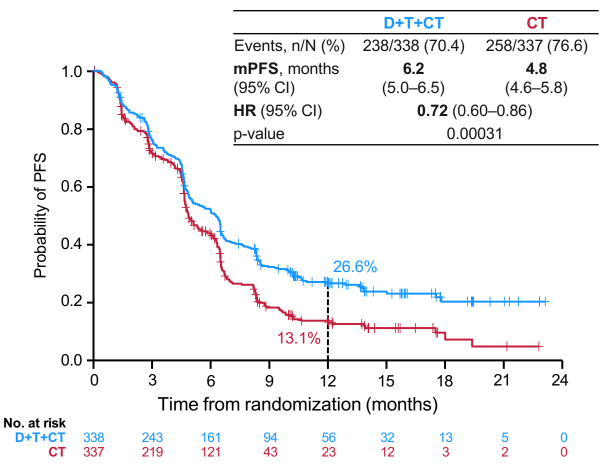


*CT options: gemcitabine + carboplatin/cisplatin (squamous), pemetrexed + carboplatin/cisplatin (non-squamous), or nab-paclitaxel + carboplatin (either histology);

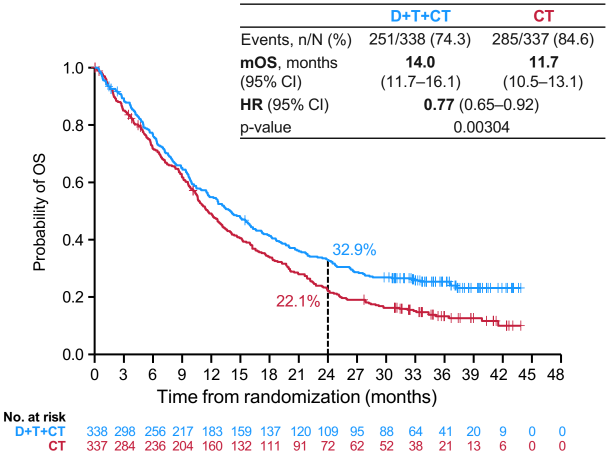
†Patients with non-squamous histology who initially received pemetrexed during first-line treatment only (if eligible); ‡Patients received an additional dose of tremelimumab post CT (5th dose)



Durvalumab + Tremelimumab + CT vs CT: PFS and OS OS







Median follow-up in censored patients at DCO: 34.9 months (range 0–44.5)



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Chemotherapy/IO Combinations

Carboplatin/Pemetrexed/Pembrolizumab [Keynote 189]

Carboplatin/Paclitaxel/Bevacizumab/Atezolizumab [IMPOWER 150]

Carboplatin/Paclitaxel or nab-paclitaxel/Pembrolizumab [Keynote 407]

• Cemiplimab/Chemotherapy [Empower Lung-3]

Durvalumab + Tremelimumab/Chemotherapy [Poseidon 3]

IO single Agent (PDL1>50%)

Pembrolizumab [Keynote 024 and 042]

Atezolizumab [IMPOWER 110]

• Cemiplimab [Empower Lung-1]

Immunotherapy combinations:

Politimumab and Nivolumab [Checkmate 227]

Ipilimumab and Nivolumab plus 2 cycles of chemotherapy [Checkmate 9LA]



ORR slightly in favor of combination chemo+IO

	KN 24	KN 42	IMPW 10 TC3/IC3	KN 407	KN 189
	(TPS > 50%)	(TPS > 50%)	(>50% and >10%)	(TPS > 50%)	(TPS > 50%)
ORR	45%	39.5%	30.7%	60.3%	61.4%
DOR	Nr (1.8-20.6 m)	20.2 m	Nr (1.8- 29.3m)	7.7 m (all patients)	11.2 m (all patients)



Adverse Events more prevalent with Chemo/IO

	KN-42		KN-24		KN-189		KN-407	
	Pembro	СТ	Pembro	СТ	Pembro + CT	СТ	Pembro + CT	СТ
All TRAE (%)	62.7%	89.9%	76.6%	90.0%	99.8%	99.0%	98.2%	97.9%
Grade 3-5 TRAE (%)	17.8%	41%	31.2%	53.3%	67.2%	65.0%	69.8%	68.2%
Discontinuation rate (any) (%)	9%	9.4%	13.6%	10.7%	27.7%	14.9%	23.4%	11.8%
Led to death	0.2%	0%	1.3%	2.0%	6.7%	5.9%	8.3%	6.4%





Outcomes of anti-PD-(L)1 therapy with or without chemotherapy (chemo) for first-line (1L) treatment of advanced non-small cell lung cancer (NSCLC) with PD-L1 score ≥50%: FDA Pooled Analysis

Oladimeji Akinboro¹, Jonathon Vallejo¹, Erica Nakajima¹, Yi Ren¹, Pallavi Mishra-Kalyani¹, Erin Larkins¹, Paz Vellanki¹, Nicole Drezner¹, Mathieu Luckson¹, Shenghui Tang¹, Martha Donoghue^{1,2}, Richard Pazdur^{1,2}, Julia A. Beaver^{1,2}, Harpreet Singh^{1,2}

¹Center for Drug Evaluation and Research, U.S. Food and Drug Administration

²Oncology Center of Excellence, U.S. Food and Drug Administration

Oladimeji Akinboro, MD, MPH









Clinical trials of first-line Chemo-IO and IO regimens included in FDA pooled analysis



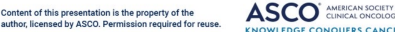
	Chemo-IO Trials	IO-only Trials		
Trial	Investigational Regimen	Trial	Investigational Regimen	
KEYNOTE-021*	Pembrolizumab + Chemo**	CheckMate 026	Nivolumab**	
KEYNOTE-189	Pembrolizumab + Chemo**	KEYNOTE-024	Pembrolizumab**	
KEYNOTE-407	Pembrolizumab + Chemo**	KEYNOTE-042	Pembrolizumab**	
IMpower150	Atezolizumab + Bevacizumab + Chemo***	IMpower110	Atezolizumab**	
IMpower130	Atezolizumab + Chemo**	CheckMate 227	Nivolumab + Ipilimumab**	
CheckMate-9LA	Nivolumab + Ipilimumab + Chemo**	EMPOWER-Lung 1	Cemiplimab**	

Abbreviations: Chemo-IO=platinum-based doublet chemotherapy immunotherapy; IO=immunotherapy.





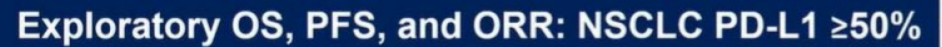




^{*} Cohort G

^{**} Control arms: Platinum-based doublet chemotherapy

^{***} Control arm in IMpower150: Bevacizumab plus platinum-based doublet chemotherapy



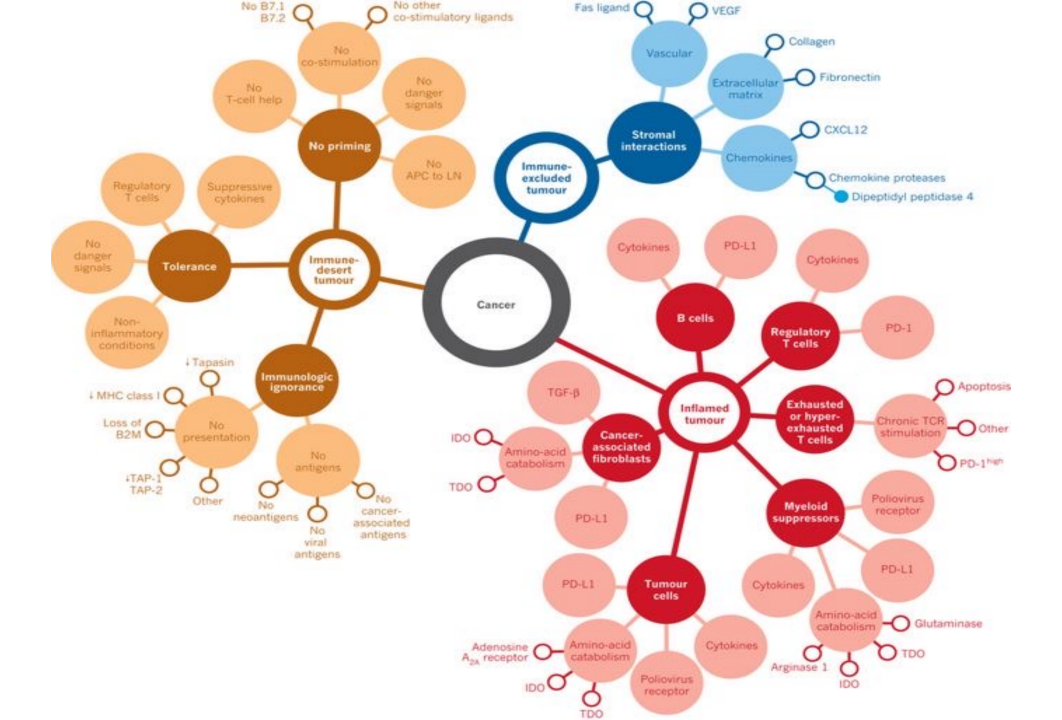


	Chemo-IO (<i>N</i> =455)		IO-alone (N=1,298)
os			
Median, months (95% CI)	25.0 (19.0, NE)		20.9 (18.5, 23.1)
HR (95% CI)		0.82 (0.62, 1.08)	
PFS			
Median, months (95% CI)	9.6 (8.4, 11.1)		7.1 (6.3, 8.3)
HR (95% CI)		0.69 (0.55, 0.87)	
ORR			
% (95% CI)	61 (56, 66)		43 (41, 46)
Odds ratio		1.2 (1.1, 1.3)	

Abbreviations: Chemo-IO=platinum-based doublet chemotherapy plus immunotherapy; CI=confidence interval; HR-hazards ratio; IO=immunotherapy; N=number; NSCLC=non-small-cell lung cancer; NE=not estimable; ORR=objective response rate; OS=overall survival; PD-L1=programmed death ligand-1; PFS=progression-free survival.









Microbiome

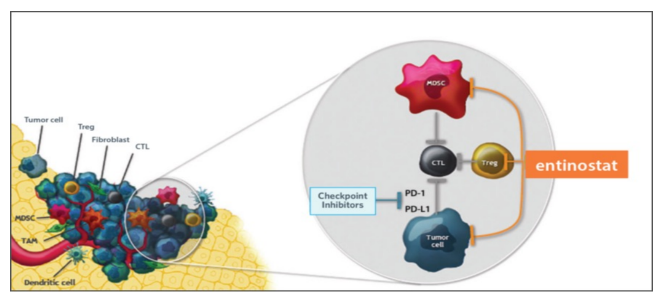
Low Benefit Conventional immunotherapy ICI alone **Patients** High Benefit High TMB **dMMR** Personalized Therapy High Benefit Tregs/MDSCs Immune Adjuvants **Predictive Biomarkers** Tumor tissue PD-L1 expression Cold tumor High Benefit TMB MMR Low Ki-67 TIL Immune Adjuvants expression Tregs/MDSCs Neoantigens Peripheral blood Low % of miRNA Exclusion Lymphocyte Tregs/MDSCs from immunotherapy (Less than 15%) Ki-67 expression Feces



CLINICAL CANCER RESEARCH | CLINICAL TRIALS: IMMUNOTHERAPY

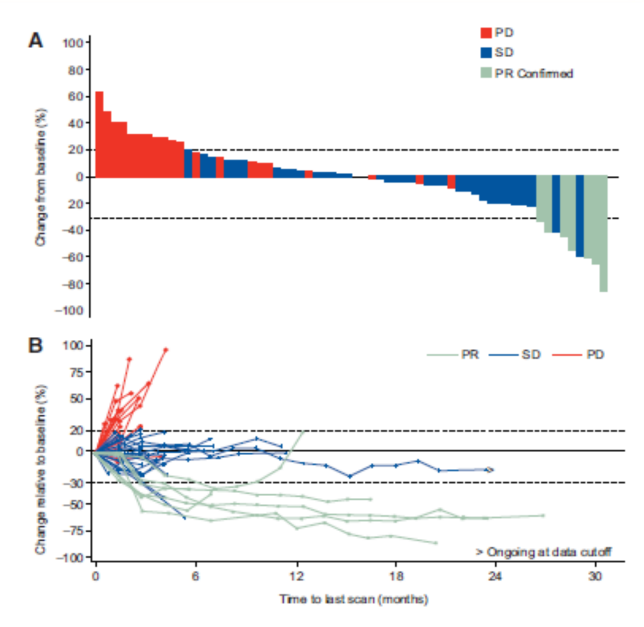
Entinostat plus Pembrolizumab in Patients with Metastatic NSCLC Previously Treated with Anti-PD-(L)1 Therapy

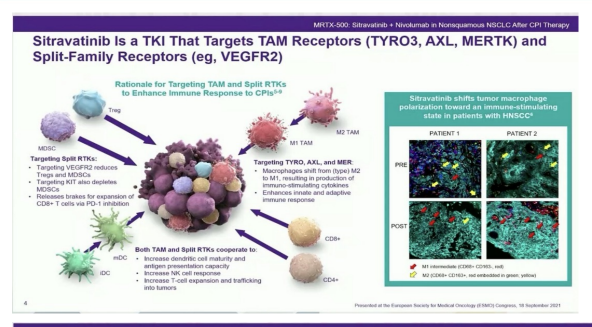
Matthew D. Hellmann¹, Pasi A. Jänne², Mateusz Opyrchal³, Navid Hafez⁴, Luis E. Raez⁵, Dmitry I. Gabrilovich⁶, Fang Wang⁶, Jane B. Trepel⁷, Min-Jung Lee⁷, Akira Yuno⁷, Sunmin Lee⁷, Susan Brouwer⁸, Serap Sankoh⁸, Lei Wang⁸, David Tamang⁸, Emmett V. Schmidt⁹, Michael L. Meyers⁸, Suresh S. Ramalingam¹⁰, Elaine Shum¹¹, and Peter Ordentlich⁸



- Entinostat (ENT) is an oral class I-selective histone deacetylase inhibitor
- ENT leads to the downregulation of immunosuppressive cell types in the tumor microenvironment
- Synergy with anti-PD1 inhibition in preclinical models
- Promising activity is shown in combination with pembrolizumab in patients with melanoma and lung cancer

- Objective response rate with ENT + PEMBRO was 10% (7 of 72, 95% CI: 4-19%)
 - Median duration of response was 5.3 months
 - An additional 50% of patients achieved disease stabilization
- Median progression-free survival = 2.8 months (95% CI: 2.1-4.1)





MRTX-500: Sitravatinib + Nivolumab in Nonsquamous NSCLC After CPI Therapy

MRTX-500: Phase 2, Open-Label Study of Sitravatinib + Nivolumab in Patients With Nonsquamous NSCLC With Prior Clinical Benefit From Checkpoint Inhibitor Therapy

Key Eligibility Criteria

- Advanced/metastatic nonsquamous NSCLC
- No actionable driver mutations
- Anti–PD-1/L1 must be the most recent line of therapy
- Prior Clinical Benefit (PCB) to CPI: CR, PR, or SD ≥12 weeks from prior CPI therapy
- No uncontrolled brain metastases
- ECOG PS 0-2

Primary Endpoint:

 Objective Response Rateb (ORR), as defined by RECIST 1.1

nivolumab

Secondary Endpoints:

- · Safety and tolerability
- · DOR
- CBR
- lity
 - PFSOS
 - 1-year survival rate

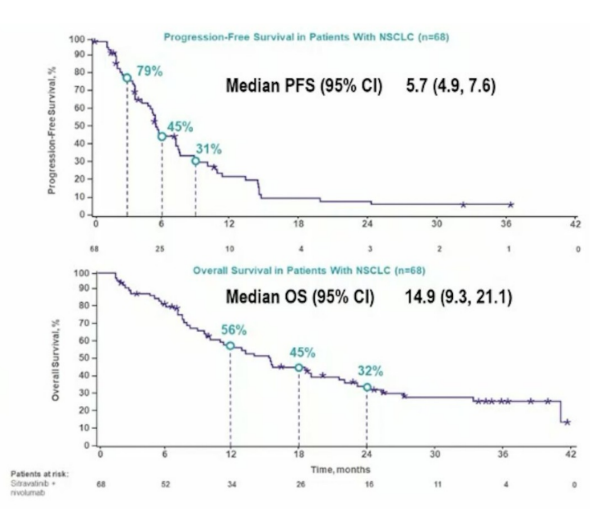
Sitravatinib 120 mg QD +

Here we report updated efficacy and safety with sitravatinib + nivolumab in the 2L or 3L setting in patients with nonsquamous NSCLC who have experienced clinical benefit on a prior CPI and subsequent disease progression

Data as of 1 June 2021

Additional cohorts included a CPI-experienced cohort that did not receive prior clinical benefit from CPI therapy (radiographic progression of disease \$12 weeks after initiation of treatment with CPI) and a CPI-naive cohort in patients that were previously treated with platinum-based chemotherapy. *Colpective response rate based on investigator assessment. Design strategies from the base formulation, involumab, 240 mg QZW or 480 mg QZW. Treatment discontinuation could be due to further treatment, study fermatable, study fermatable, or death.

Presented at the European Society for Medical Oncology (ESMO) Congress, 18 September 2021



Ticiana Leal. ESMO 2021.

CONQUERING THORACIC CANCERS WORLDWIDE

STK11/LKB1, KRAS mutations and immunerelated adverse events as predictors of response to immunotherapy in lung cancer

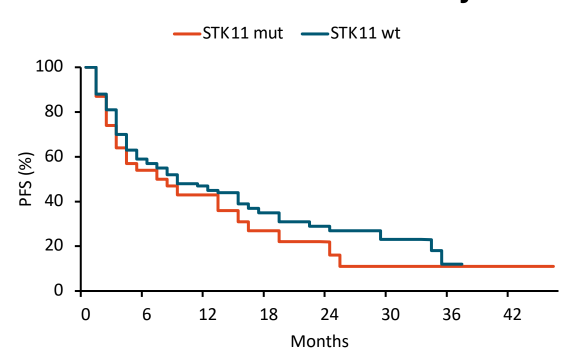
Luis E. Raez, MD¹; Richie Uba, PharmD²,³; Aaron North, PharmD²,³; Katerine Dumais, PharmD, MPH¹; Hermán W. Powery II, PharmD, BCOP¹; Gelenis Domingo, MD¹; Brian Hunis, MD¹; Paola Izquierdo, ARNP¹; Frank Gentile, PharmD, BCOP¹

¹Memorial Cancer Institute, Pembroke Pines, FL; ²Florida A&M University, Davie, FL; ³Memorial Regional Hospital, Hollywood, FL

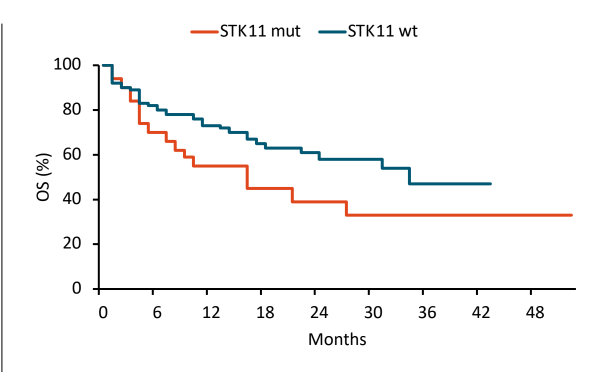
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CONQUERING THORACIC CANCERS WORLDWIDE

Results: PFS and OS by STK11 Status



	STK11 wt n = 96	STK11 mut n = 31	HR (95% CI)	р
mPFS	6.3m	5.6m	1.35 (0.76-2.1)	0.35
12-m PFS	45%	43%	-	0.85



	STK11 wt n = 96	STK11 mut n = 31	HR (95% CI)	р
mOS	12.1m	8.6m	1.7 (1.0-3.6)	0.03
12-m OS	73%	55%	-	0.03









Overall survival from a phase II randomized study of ramucirumab plus pembrolizumab versus standard of care for advanced non-small cell lung cancer previously treated with immunotherapy—Lung-MAP non-matched sub-study S1800A

Karen L. Reckamp, M.D.¹, Mary W. Redman, PhD², Konstantin H. Dragnev, M.D.³, Liza Villaruz, M.D.⁴, Bryan Faller, MD⁵; Tareq Al Baghdadi, MD⁶, Susan Hines, MD⁷, Lu Qian, M.S.², Katherine Minichiello, M.S.², David R. Gandara, M.D.⁸, Karen Kelly, MD⁸, Roy S. Herbst, M.D., Ph.D.⁹

¹Cedars-Sinai Medical Center, Los Angeles, CA; ²SWOG Statistics and Data Management Center & Fred Hutchinson Cancer Research Center, Seattle, WA; ³Dartmouth-Hitchcock Norris Cotton Cancer Center, Lebanon, NH/Alliance for Clinical Trials in Cancer; ⁴University of Pittsburgh Medical Center (UPMC) Hillman Cancer Center; ⁵Missouri Baptist Medical Center, St. Louis, MO/Heartland NCORP; ⁶IHA Hematology Oncology Consultants-Ann Arbor/Michigan CRC NCORP; ⁷Novant Health Cancer Institute - Mount Airy/Southeast Clinical Oncology Research Consortium NCORP); ⁸UC Davis Comprehensive Cancer Center, Sacramento, CA; ⁹Yale University, New Haven, CT







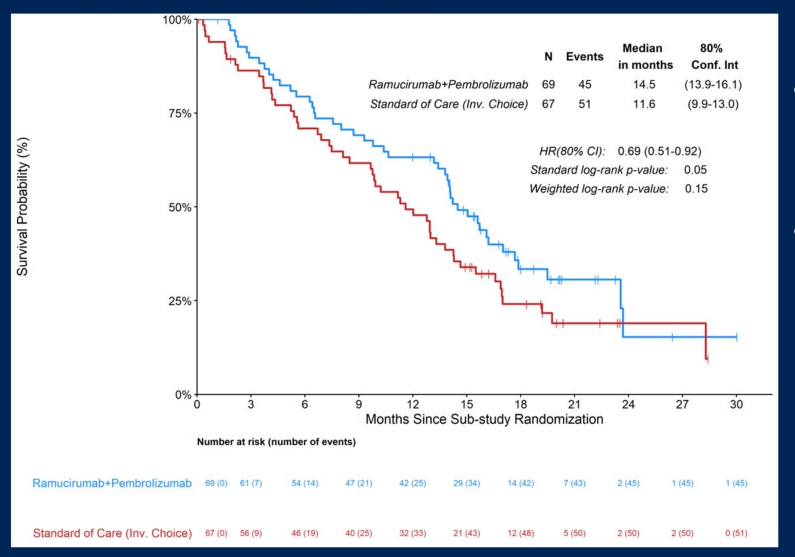








Overall survival



 Median OS for RP 14.5 months v. SOC 11.6 months

HR= 0.69; SLR p-value 0.05

Standard of care therapy received:

- Docetaxel + Ramucirumab (n = 45)
- Docetaxel (n = 3)
- Gemcitabine (n = 12)
- Pemetrexed (n = 1)
- No treatment (n = 6)







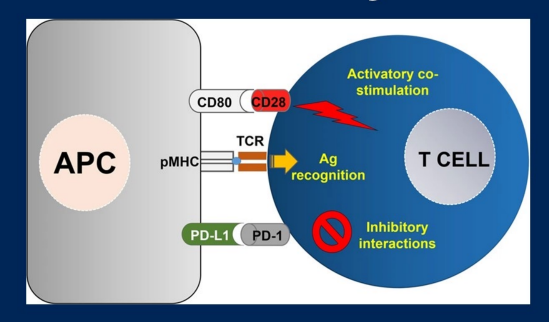


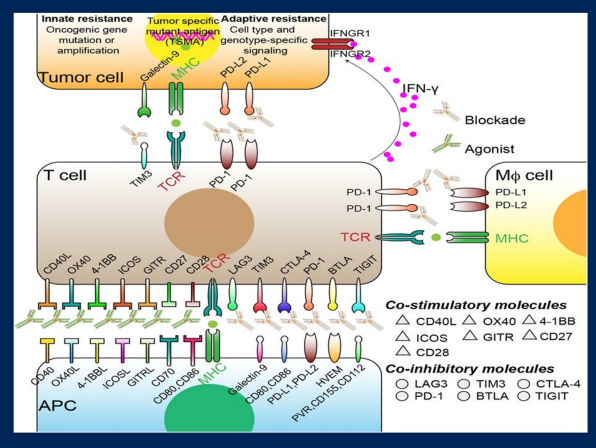


Immunotherapy resistance in NSCLC

- Deacetylase Inhibitors (etinostat, vorinostat)
- Vaccines (Heat Shock Protein gp96)
- STK11/LKB1 and KEAP
- Cytokines (IL-10, IL-15)
- Adenosine pathway
- Combination checkpoints (IDO, TIM, LAG3, TIGIT)
- Adoptive T cell therapies (TILs, TILs + anti PD(L)-1, TCRs)
- B-catenin pathway

Co-Stimulatory and Co-Inhibitory Interactions





Escors, et al Signal Transduct Target Ther 2018

Li Cellular and Molecular Immunology 2018

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Thanks







