

Immunotherapy for the Treatment of Skin Cancers

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Disclosures

- Consultant: Array
- Research funding (paid to institution): Bristol Myers Squibb, Dynavax, Immunocore, Merck
- I will be discussing non-FDA approved indications during my presentation.





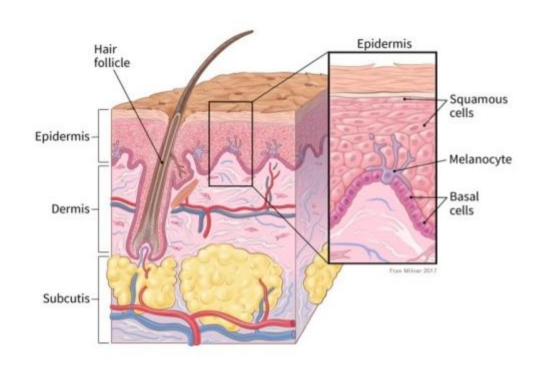






Background

- Skin cancer is the most common type of cancer
- Three most common types of skin cancers:
 - Basal cell carcinoma
 - Squamous cell carcinoma
 - Melanoma
- Melanoma was one of the foundational disease states for testing immunotherapies













Approved cytokines in melanoma

Drug	Indication	Dose
High-dose interferon alfa-2b	Adjuvant – high risk for systemic recurrence	Induction: 20m IU/m ² IV 5x/wk for 4 wks Maintenance: 10m IU/m ² s.c. 3x/wk for 48 wks
Interleukin-2 (Aldesleukin)	Stage IV	600k IU/kg/dose Q8hr, up to 14 doses; 9 days of rest; can repeat up to 28 doses per course
Pegylated Interferon alfa-2b (Sylatron)	Adjuvant – microscopic or gross nodal involvement	6 mcg/kg/wk s.c. for 8 doses, then 3 mcg/kg/wk s.c. for up to 5 years











Approved checkpoint inhibitors in melanoma

Drug	Approved	Indication	Dose
Ipilimumab	2011	Unresectable/Metastatic melanoma: newly diagnosed or after progression	3 mg/kg Q3W for 4 doses
	2015	Adjuvant therapy in stage III melanoma after complete resection	10 mg/kg Q3W for 4 doses, then 10 mg/kg Q12W for 3 years
	2017	Unresectable/Metastatic melanoma: newly diagnosed or after progression, all patients ≥ 12 yr	3 mg/kg Q3W for 4 doses





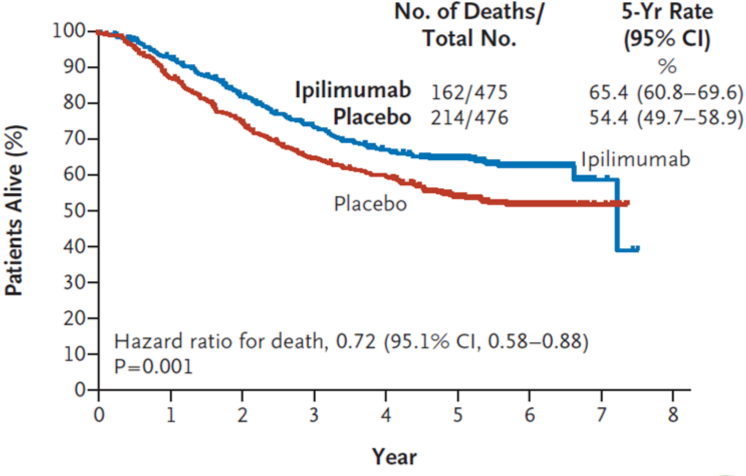






Adjuvant Ipilimumab in High-Risk Stage III Melanoma

- EORTC 18071 phase III trial
 - NCT00636168
 - Adjuvant ipilimumab vs placebo
 - Ipilimumab 10mg/kg Q3W for four doses, then every 12 weeks for up to 3 years











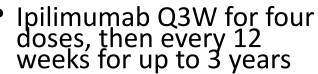


Adjuvant Ipilimumab in High-Risk Stage III Melanoma

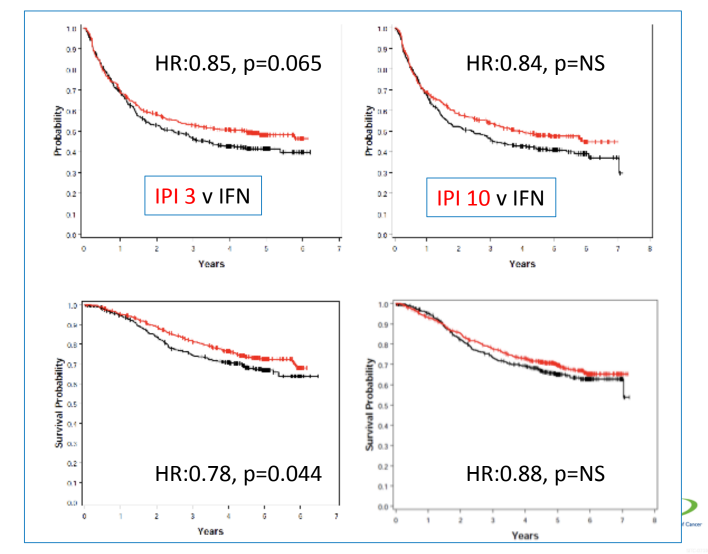
- ECOG 1609
 - NCT01274338
 - Adjuvant interferon (IFN) vs ipilimumab 3 mg/kg (IPI 3) vs ipilimumab 10 mg/kg (IPI 10)

 - IPI 3 "better than IFN", IPI 10 "not better than IFN"
 - IPI3 better tolerated than IPI 10

RFS



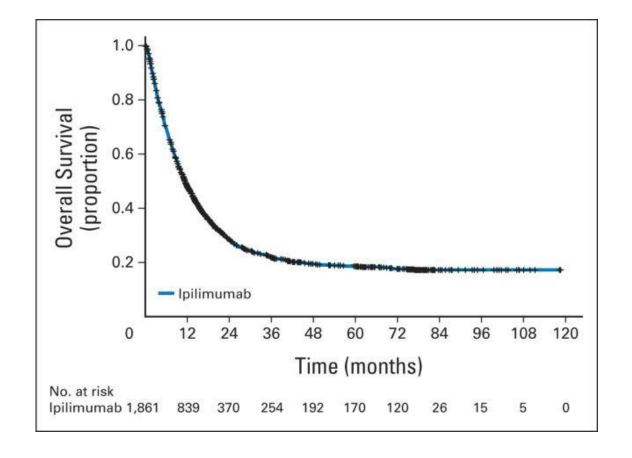
OS





Ipilimumab in Stage III/IV Melanoma

- Pooled OS data from 10 phase II/III trials
 - Previously treated (n = 1,257) or treatmentnaïve (n = 604)
 - Ipilimumab 3 mg/kg (n = 965) or 10 mg/kg (n = 706)













Approved checkpoint inhibitors in melanoma

Drug	Approved	Indication	Dose	
	2014	Advanced/unresectable melanoma with progression after other therapy	200 mg Q3W*	
Pembrolizumab	2015	1 st line unresectable/metastatic melanoma	200 mg Q3W*	
	2019	Adjuvant therapy of melanoma following complete resection	200 mg Q3W	
*Original approvals were 2 mg/kg Q3W – updated to flat dosing regimen				





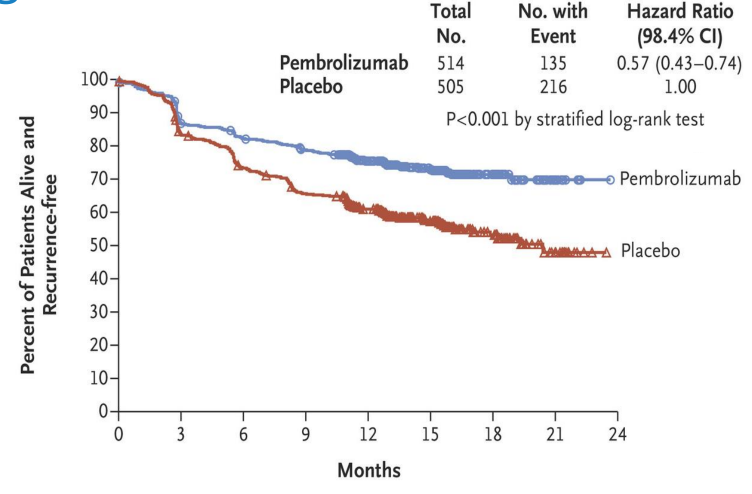






Adjuvant Pembrolizumab in High-Risk Stage III Melanoma

- EORTC 1325/KEYNOTE-054 phase III trial
 - NCT02362594
 - Adjuvant pembrolizumab vs placebo
 - Pembrolizumab 200mg
 Q3W for up to 1 year
 (~18 total doses)







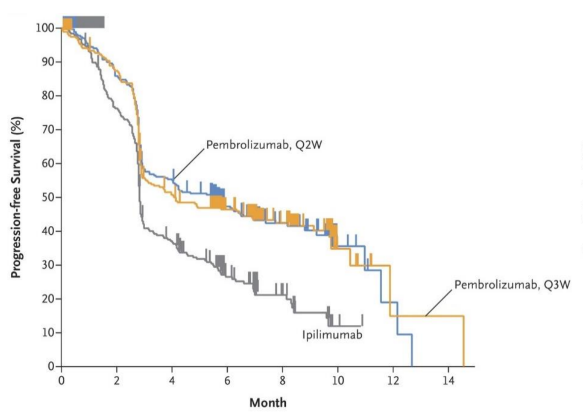


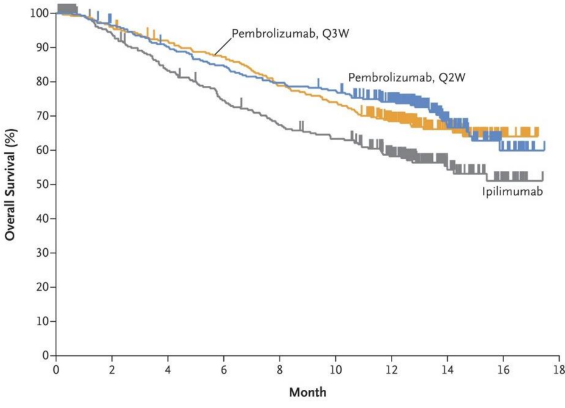




Pembrolizumab in Stage III/IV Melanoma

Phase III KEYNOTE-006 Trial















Approved checkpoint inhibitors in melanoma

Drug	Approved	Approved Indication		
Nivolumab	2014	Unresectable/metastatic melanoma with progression after other therapy	240 mg Q2W or 480 mg Q4W*	
	2017	Adjuvant treatment of melanoma after complete resection	240 mg Q2W or 480 mg Q4W	
*Original approval was 3 mg/kg Q2W, updated to flat dosing regimen				





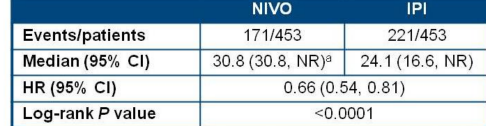


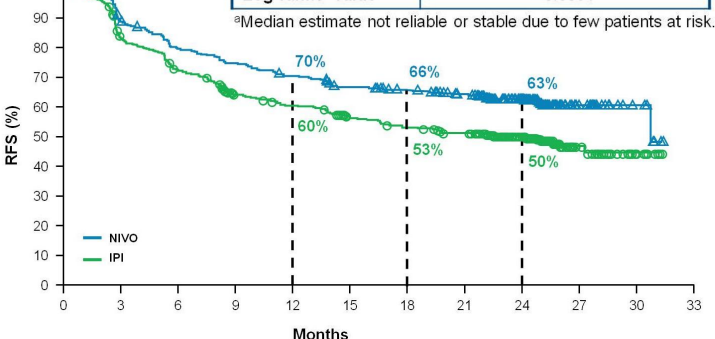




Adjuvant Nivolumab vs Ipilimumab in High-Risk Stage III Melanoma

- CheckMate 238 phase III trial
 - NCT02388906
 - Ipilimumab 10mg/kg Q3W for four doses, then every 3 months for up to 1 year
 - Nivolumab 3mg/kg
 Q2W for four doses,
 then every 3 months for
 up to 1 year















Approved checkpoint inhibitors in melanoma

Drug	Approved	Indication	Dose
Nivolumah ı Inilimumah	2015	BRAF V600 WT unresectable/metastatic melanoma	1 mg/kg nivolumab + 3 mg/kg ipilimumab Q3W for 4 doses, then nivolumab 240 mg Q2W or 480 mg Q4W
Nivolumab + Ipilimumab	2016	BRAF V600 WT or mutant unresectable/metastatic melanoma	1 mg/kg nivolumab + 3 mg/kg ipilimumab Q3W for 4 doses, then nivolumab 240 mg Q2W or 480 mg Q4W





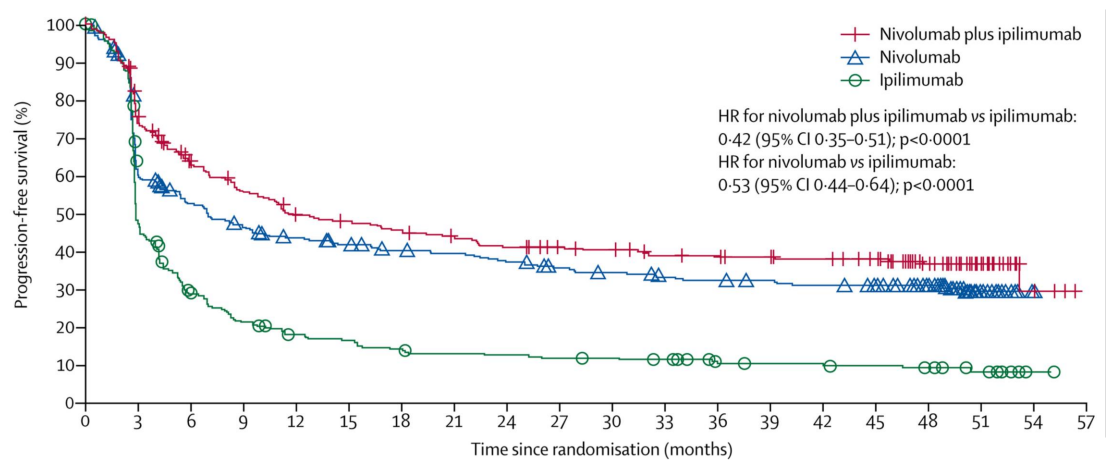






Combination Ipilimumab + Nivolumab in Stage III/IV Melanoma

Phase III CheckMate 067 Trial







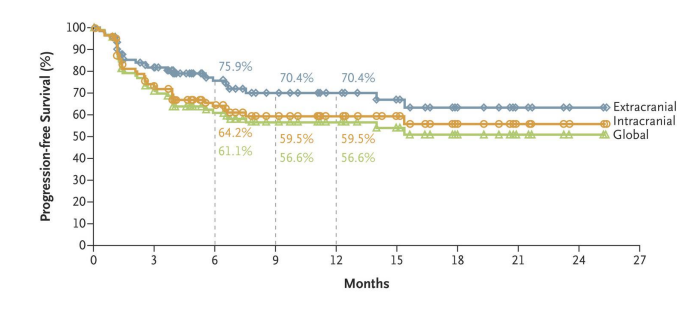






Combination Ipilimumab + Nivolumab for Patients with Asymptomatic Brain Metastases

Variable	Intracranial (N = 94)	Extracranial (N = 94)	Global (N = 94)
Best overall response — no. (%)*			
Complete response	24 (26)	7 (7)	8 (9)
Partial response	28 (30)	40 (43)	40 (43)
Stable disease for ≥6 mo	2 (2)	6 (6)	5 (5)
Progressive disease	31 (33)	28 (30)	33 (35)
Could not be evaluated†	9 (10)	13 (14)	8 (9)
Objective response‡			
No. of patients	52	47	48
Percent of patients (95% CI)	55 (45–66)	50 (40–60)	51 (40–62)
Clinical benefit§			
No. of patients	54	53	53
Percent of patients (95% CI)	57 (47–68)	56 (46–67)	56 (46–67)





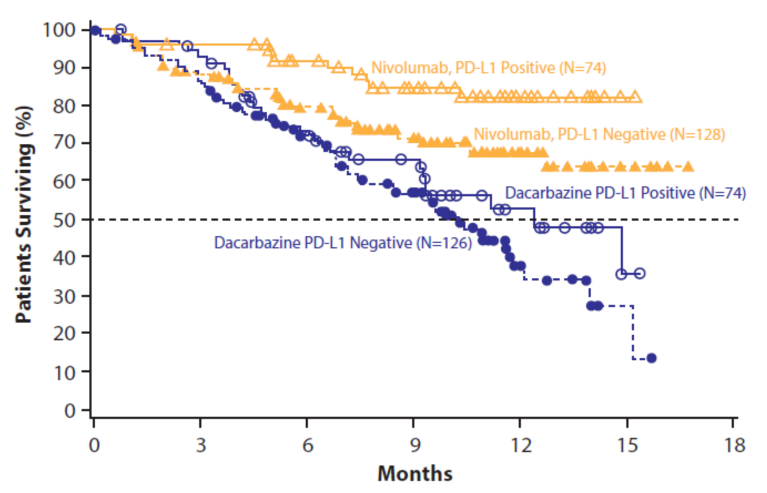








Importance of Tumor PD-L1 Status with Anti-PD-1 Monotherapy



	Patients Who Died n/N	Median Survival mo (95% CI)
Nivolumab PD-L1 Positive	11/74	N.R.
Nivolumab PD-L1 Negative	37/128	N.R.
Dacarbazine PD-L1 Positive	29/74	12.4 (9.2-N.R.)
Dacarbazine PD-L1 Negative	64/126	10.2 (7.6–11.8)



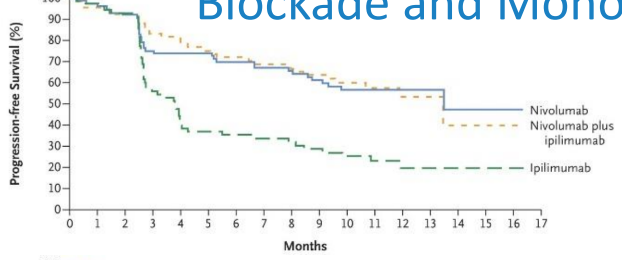




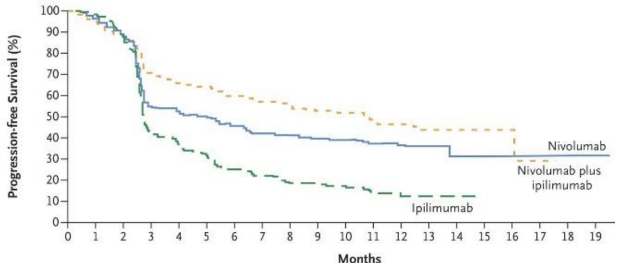




Importance of Tumor PD-L1 Status between Combination Checkpoint Blockade and Monotherapy



Tumor PD-L1 Positive Patients



Tumor PD-L1 Negative Patients



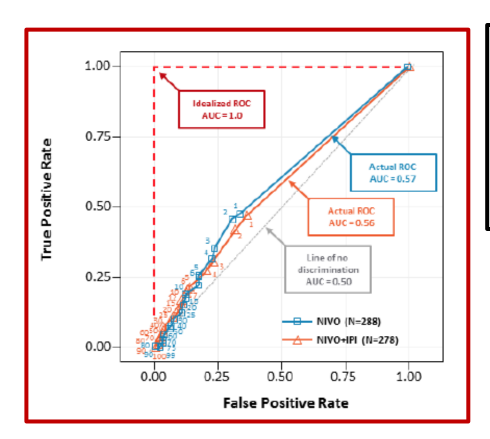








The use of PD-L1 status to predict overall survival is poor with single-agent PD-1 or combined ipi/nivo...



PDL-1 (%)	≥1	< 1	≥ 5	< 5	> 10	< 10
Ipilimumab	19%	18%	21%	17%	20%	18%
Nivolumab	54%	35%	58%	42%	58%	44%
lpi/Nivo	65%	54%	72%	56%	85%	55%

...but, PD-L1 status predicts higher response rate with combo at every PD-L1 expression cut-off











In development: Neoadjuvant immunotherapy in advanced melanoma

Trial	Regimen	N	pCR (%)	med RFS (mo)	med FU (mo)
Amaria Lancet Oncol 2018	Dab/Tram	21	58	19.7	18.6
Long Lancet Oncol 2019	Dab/Tram	35	49	23.0	27.0
Blank Nat Med 2018	lpi+nivo	10	33	NR	32
Amaria Nat Med 2018	Nivo Ipi+nivo	12 11	25 45	NR NR	20
Huang Nat Med 2019	Pembro	30	19	NR	18
Rozeman Lancet Oncol 2019	lpi+nivo	86	57	NR	8.3





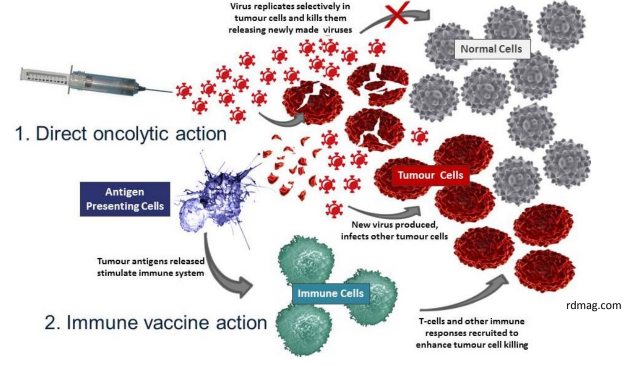






Approved oncolytic virus in

melanoma



Drug	Approved	Indication	Dose
imogene epvec (T-Vec)	2015	Local treatment of unresectable cutaneous, subcutaneous, and nodal lesions in recurrent melanoma after surgery	Intralesional injection: ≤4 mL at 10 ⁶ PFU/mL starting; 10 ⁸ PFU/mL subsequent







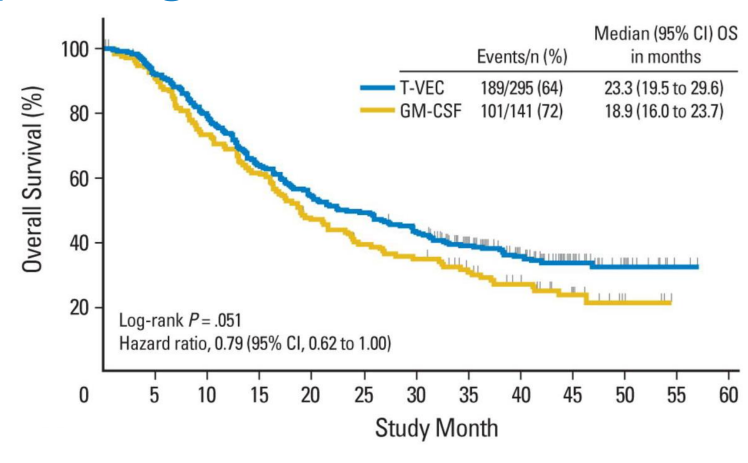




Talimogene laherparepvec (T-VEC) in Stage III/IV Melanoma

Phase III OPTiM Trial

- Oncolytic, geneticallyengineered herpes virus
- Intralesional T-VEC
 106 pfu/mL,
 108 pfu/mL 3 weeks
 after initial dose, then
 Q2W
- Subcutaneous GM-CSF













Approved checkpoint inhibitors in other skin cancers

Drug	Approved	Indication	Dose
Avelumab	2017	Patients >12 yr with metastatic Merkel cell carcinoma	800 mg Q2W + premedication (first 4 cycles)
Pembrolizumab	2018	Adult/pediatric with recurrent advanced/metastatic Merkel cell carcinoma	Adults: 200 mg Q3W Pediatric: 2 mg/kg (up to 200 mg) Q3W
Cemiplimab-rwlc	2018	Metastatic cutaneous squamous cell carcinoma, not candidate for curative therapies	350 mg Q3W





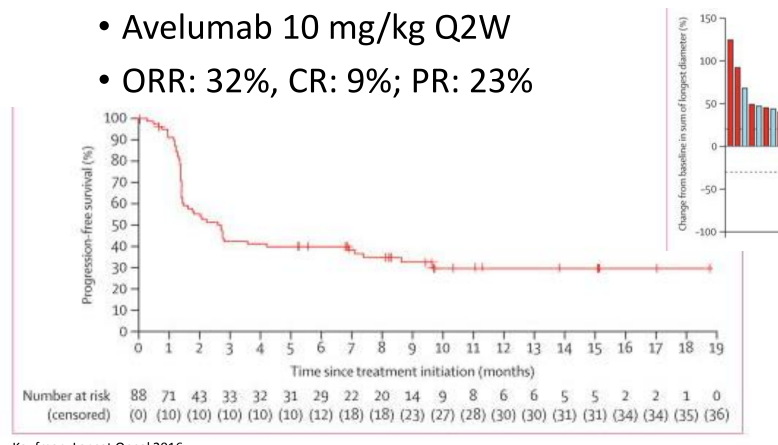






Avelumab in 2nd-line metastatic Merkel Cell carcinoma

• 1st FDA-approved treatment for this status







One previous line of any systemic therapy (n=39)
 Two or more previous lines of any systemic therapy (n=26)

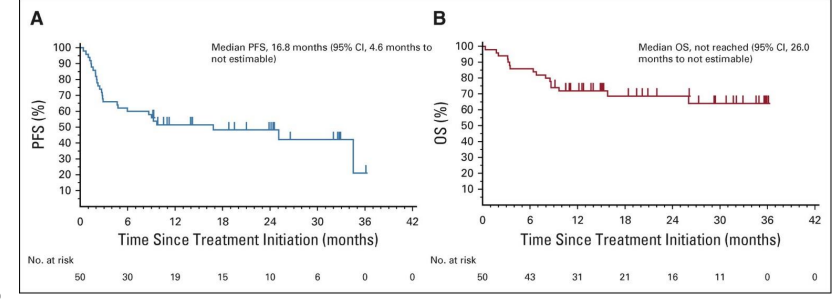






Pembrolizumab in 1st-line advanced Merkel Cell Carcinoma

- KEYNOTE-017
- Pembrolizumab 2 mg/kg Q3W up to 2 years
- mPFS: 16.8 months (compared to 90 days for chemo)
- 24-month OS: 68.7%





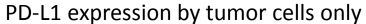








Pembrolizumab in 1st-line advanced Merkel Cell Carcinoma

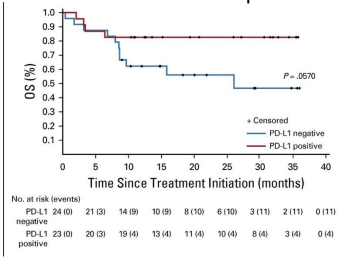


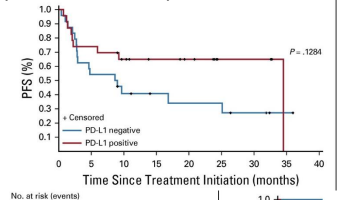
negative

positive

PD-L1 23 (0)

17 (6)

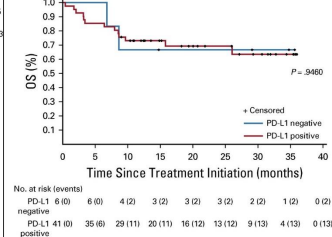


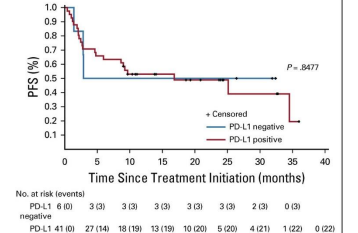


13 (11) 8 (14) 6 (14) 5 (15)

13 (8)











positive

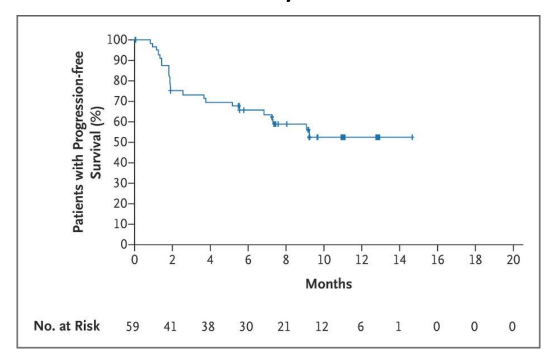


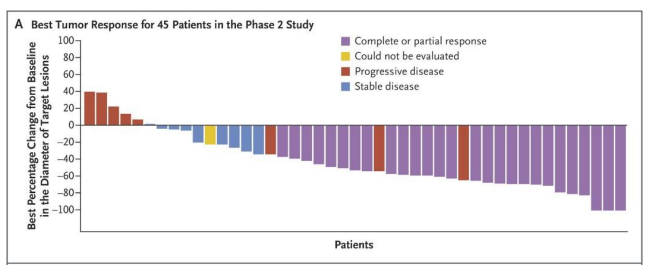




Cemiplimab in advanced/metastatic cutaneous squamous-cell carcinoma

- Cemiplimab 3mg/kg Q2W
- 47% response rate in metastatic patients
- 60% of locally advanced had objective response





Migden, NEJM 2018.





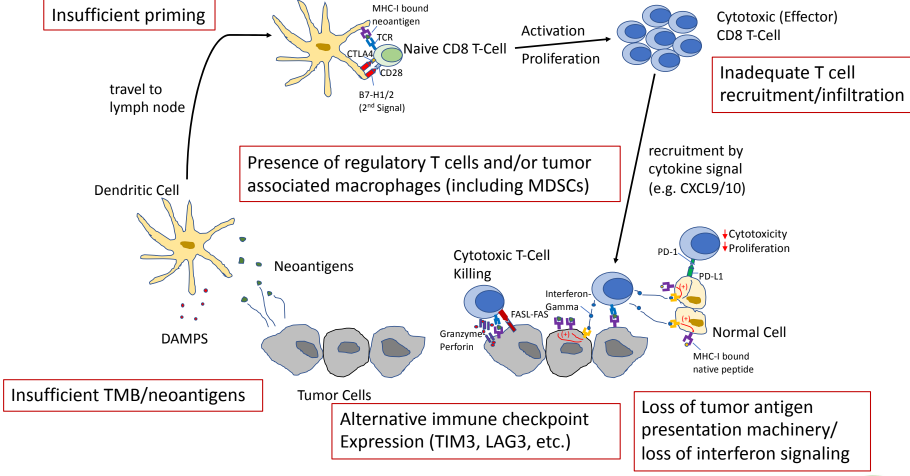






Developmental Immunotherapeutic Strategies for Melanoma

How does immune checkpoint inhibitor therapy fail?









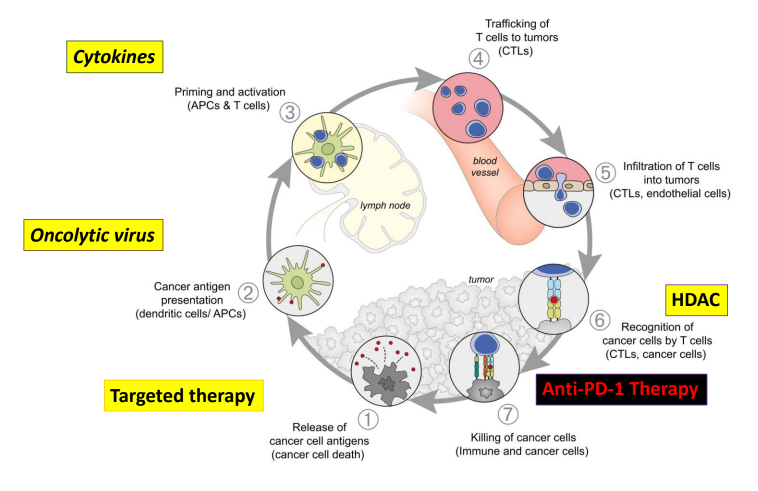




Developmental Immunotherapeutic Strategies for Melanoma

How do we overcome resistance?

Combination therapy







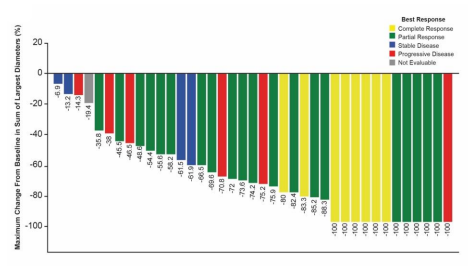




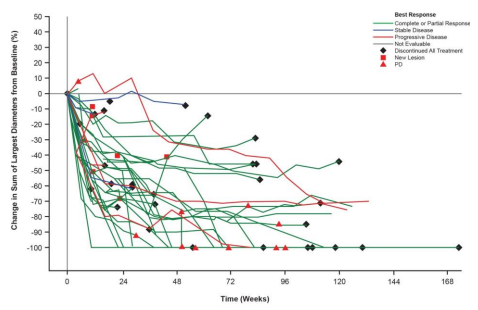


In development: Combined IO with **BRAF** targeted therapy

- Cobimetinib + vemurafenib + atezolizumab
- ORR: 71.8%
- Median duration of response: 17.4 mo









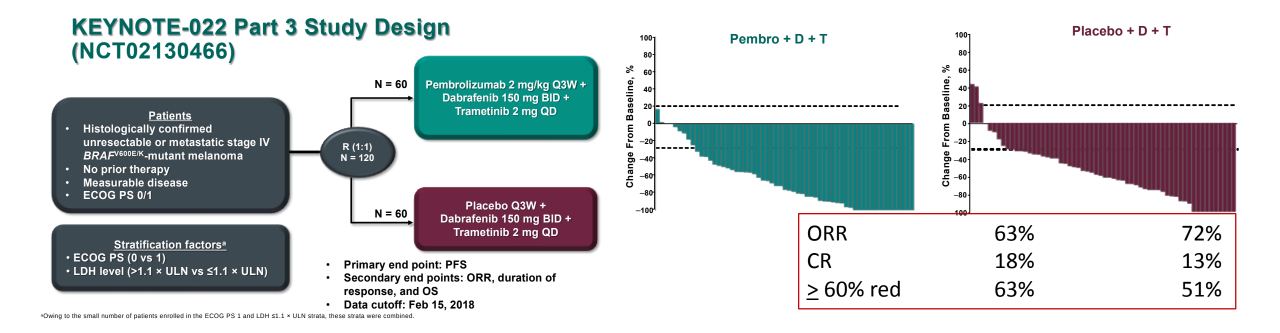








In development: Combined IO with BRAF targeted therapy





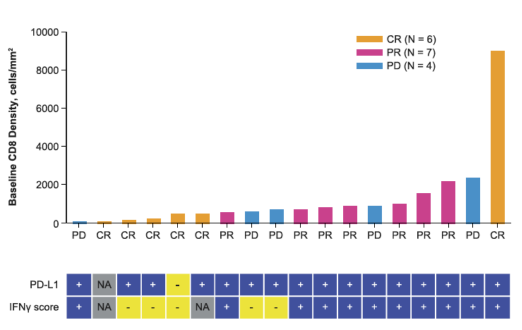




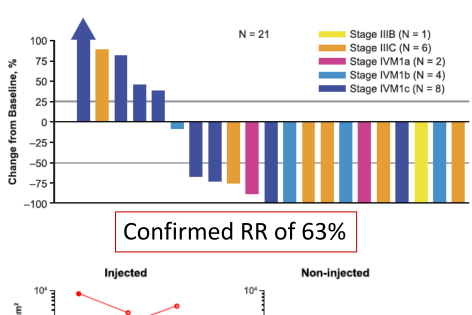


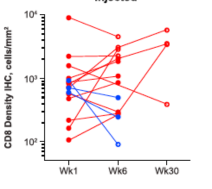


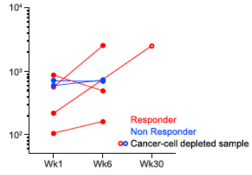
In development: Combined IO with Oncolytic Virus



Phase I: Pembrolizumab + TVEC







Ribas et al Cell 2017







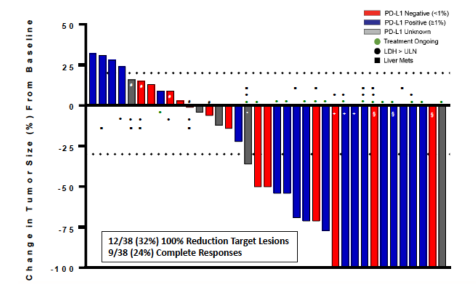




In development: Combined IO with IL-2 (NKTR-214)

Efficacy (response rate)
data from nonrandomized cohorts of
urothelial bladder cancer,
renal cell carcinoma, and
melanoma looks
promising

Stage IV IO-Naïve 1L Melanoma Cohort at RP2D Best Overall Response by Independent Radiology



1L Melanoma (n=38 Efficacy Evaluable)	Overall Response Rate
Confirmed ORR (CR+PR)	20 (53%)
CR	9 (24%)
DCR (CR+PR+SD)	29 (76%)
PD-L1 negative (n=14)	6 (43%)
PD-L1 positive (n=19)	13 (68%)
PD-L1 unknown (n=5)	1 (20%)
LDH > ULN (n=11)	5 (45%)
Liver metastases (n=10)	5 (50%)

High level of concordance in ORR between independent central radiology (53%) and investigator-assessed 19/38 (50%).







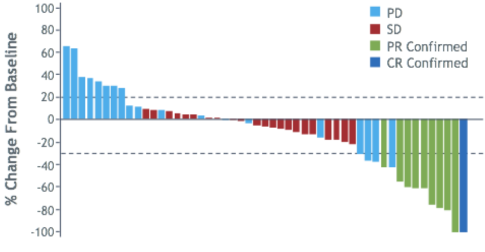


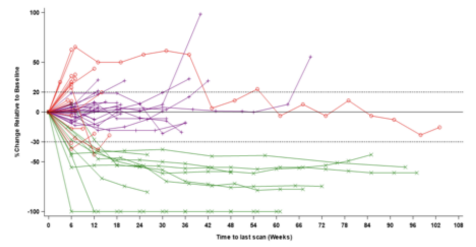


In development: Combined IO with

HDAC inhibitor

- Entinostat + pembrolizumab
- 19% ORR (1 CR, 9 PR)
- Median duration of response: 13 mo
- 9 additional patients with SD for >6 mo















Conclusions

- Melanoma was one of the foundational disease states for testing immunotherapies
- Avelumab and pembrolizumab are now approved for Merkel cell carcinoma, and cemiplimab is approved for cutaneous squamous cell carcinoma
- Combination immunotherapies may lead to higher response rates and more durable responses











Additional Resources



Sullivan et al. Journal for ImmunoTherapy of Cancer (2018) 6:44 https://doi.org/10.1186/s40425-018-0362-6

Journal for ImmunoTherapy of Cancer

POSITION ARTICLE AND GUIDELINES

Open Access



An update on the Society for Immunotherapy of Cancer consensus statement on tumor immunotherapy for the treatment of cutaneous melanoma: version 2.0

Ryan J. Sullivan¹, Michael B. Atkins², John M. Kirkwood³, Sanjiv S. Agarwala⁴, Joseph I. Clark⁵, Marc S. Ernstoff⁶, Leslie Fecher⁷, Thomas F. Gajewski⁸, Brian Gastman⁹, David H. Lawson¹⁰, Jose Lutzky¹¹, David F. McDermott¹², Kim A. Margolin¹³, Janice M. Mehnert¹⁴, Anna C. Pavlick¹⁵, Jon M. Richards¹⁶, Krista M. Rubin¹, William Sharfman¹⁷, Steven Silverstein¹⁸, Craig L. Slingluff Jr¹⁹, Vernon K. Sondak²⁰, Ahmad A. Tarhini²¹, John A. Thompson²², Walter J. Urba²³, Richard L. White²⁴, Eric D. Whitman²⁵, F. Stephen Hodi²⁶ and Howard L. Kaufman^{1*}











Case Studies











- 69 yo female
- 1970s: melanoma right forearm, with "lymph nodes removed"
- 2014:
 - left back lesion, Clark IV, Breslow 2.5mm, mitotic rate 8, ulceration present, absence of intraepidermal component-metastasis cannot be ruled out
 - WLE and SLNBx of left axilla (2/2 nodes positive; largest nodal metastasis at least 6mm, no extranodal extension)
- Adjuvant pegylated IFN
 - Held after 4 months



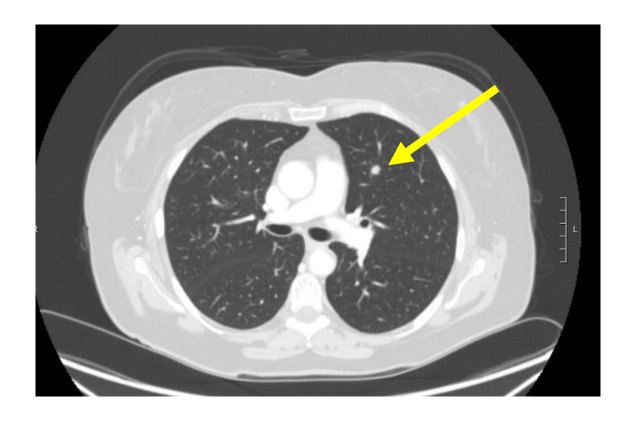


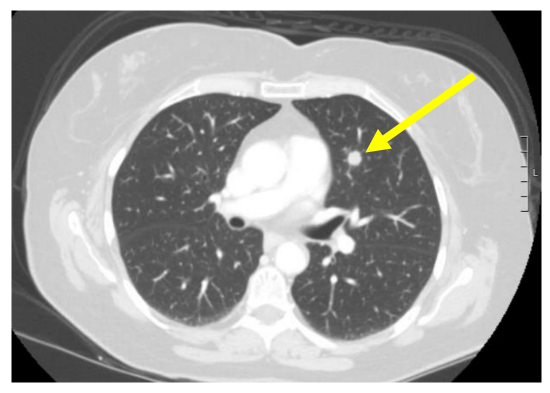






• 2018





3 months later











- Lung biopsy: + for melanoma
- What would you do as next step?
 - A. Surgery
 - B. Anti-PD-1 therapy
 - C. Anti-PD-1/anti-CTLA-4 therapy
 - D. BRAF/MEK inhibitor therapy











- Best option would be systemic therapy
 - No OS or prospective data with surgical resection
 - Best front-line therapy for melanoma debated
 - Immunotherapy should include an anti-PD-1 agent
 - Higher rates of toxicity with dual checkpoint
 - BRAF/MEKi therapies: no molecular information on this patient yet; an option for approximately 50% of patients with a BRAF^{V600E} mutation; concerns about durability of responses



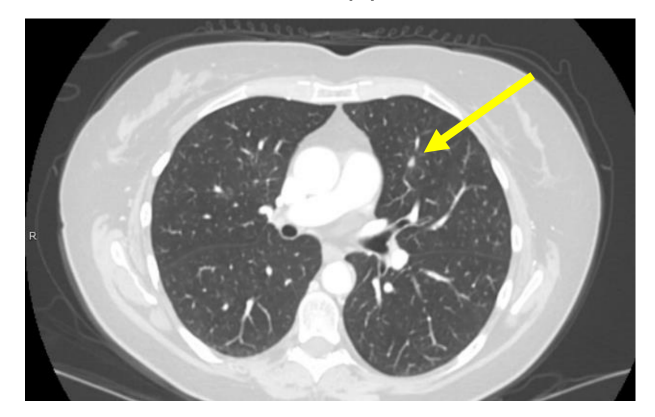








- Initiated on nivolumab monotherapy
- 2019:













• After approximately 1 year on therapy:















- Next steps?
- Options/decisions:
 - Disease progression or immune mediated toxicity
 - Continuation of therapy
 - Diagnostic considerations
 - Biopsy: Alveolar lung tissue with reactive pneumocytes and mild chronic inflammation; No evidence of malignancy











- Additional history:
- Patient reported history of e-cigarette use
- Recent change in e-cigarette liquid: made in a local store
- Current status:
 - Holding further therapy
 - Referral to pulmonology





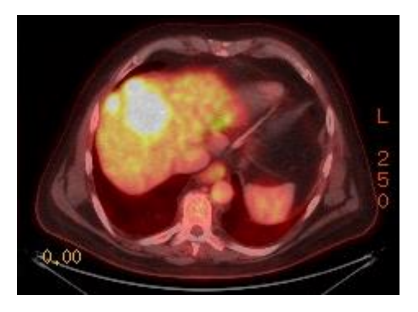






- 70 yo diagnosed with stage III melanoma 2010; received IFN alfa completed in 2011
- 2015: Progressive fatigue, malaise, SOB















- Biopsy of lung mass is performed and is positive for metastatic melanoma
- Brain MRI negative
- Molecular studies pending
- Next steps (show of hands):
- A. Wait for molecular studies before making treatment decision
- B. Treat with immunotherapy











- Treatment of patients with metastatic melanoma and high symptom burden and unknown BRAF status is challenging
- Prolonged time for BRAF results

Agent	High response rate	Quick response	Response durability	Toxicity
Anti-PD-1 monotherapy	0	0	0	0
Dual checkpoint inhibition	0		0	0
BRAF + MEK inhibition	0	0	0	0











- Patient initiated therapy with ipilimumab/nivolumab
- Presents for C3 with diarrhea
 - Abdominal pain/cramping
 - 8-10 bowel movements a day
- Next steps (show of hands):
 - A. Supportive care with loperamide
 - B. Initiation of prednisone 1mg/kg day
 - C. Infliximab
 - **D.**Colonoscopy











Option B is best choice

- Steroids are generally indicated as initial management for grade 3 colitis
- Routine colonoscopy no longer indicated
 - Consider when not improving
 - Prolonged course of steroids
 - Concern for underlying malignancy
- Infliximab often reserved for steroid refractory cases











- Patient received high dose steroids
 - Initially improved
 - Difficulty tapering
 - Infliximab X 1 with resolution



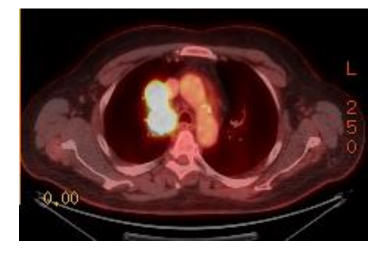




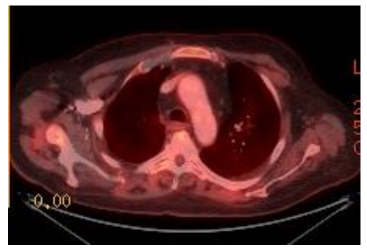


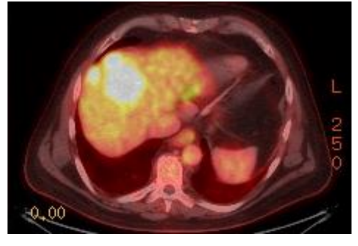


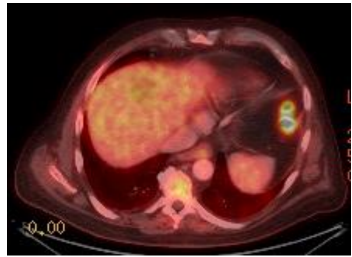
Pre-treatment



Post-treatment (no additional therapy)

















Thank You







