



Immunotherapy for the Treatment of Skin Cancers

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- Consulting: Amgen, Genmab, Xencor, BMS, Regeneron
- Contracted Research: Xencor, Astellas, Kite Pharma, Vedanta, Merck, Boehringer Ingelheim
- 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 tumor types for which immunotherapy was tested and provided proof of concept







- Melanoma
 - Front-line treatment
 - Second-line or later
 - Adjuvant and neoadjuvant settings
- Merkel cell carcinoma
- Squamous cell and basal cell carcinoma
- Future areas of research





Checkpoint Inhibitor Immunotherapy



Sharma, Hu-Lieskovan, Wargo, Ribas. Cell, 2017 Hu-Lieskovan and Ribas. Cancer Journal. 2017 AAEM AMERICAN ACADEMY OF EMERGENCY MEDICINE

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Immunotherapy Transformed Cancer Treatment

All cancer patients have the potential to respond to immunotherapy



Larkin et al. N Engl J Med 2019; 381:1535-1546



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Immunotherapy treatment options for metastatic melanoma

Treatment	Indication	Dose
Ipilimumab	Unresectable/Metastatic melanoma: newly diagnosed or after progression, all patients ≥ 12 yr	3 mg/kg Q3W for 4 doses
Pembrolizumab	Unresectable/metastatic melanoma	200 mg Q3W or 400 mg Q6W
Nivolumab	Unresectable/metastatic melanoma	240 mg Q2W or 480 mg Q4W
Nivolumab + ipilimumab	Unresectable/metastatic melanoma	1 mg/kg nivo followed by 3 mg/kg ipi Q3W, Maintenance: nivolumab 240 mg Q2W or 480 mg Q4W
Atezolizumab + cobimetinib + vemurafenib	BRAF V600 mutation-positive unresectable/metastatic melanoma	28-day cycle of cobi/vem, then atezolizumab 840 mg every 2 weeks with cobimetinib 60 mg orally once daily (21 days on/7 days off) and vemurafenib 720 mg orally twice daily
Talimogene laherparepvec (T-Vec)	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





Trials leading to initial approvals

Trial	Treatment arms	n	Patient selection criteria	ORR	Median OS (months)	Median PFS (months)
	Ipilimumab + gp100	403	Pretreated	5.7%	10.0	2.76
NCT00094653	Ipilimumab	137	advanced melanoma	10.9%	10.1	2.86
	Gp100	136	melanoma	1.5%	6.4	2.76
	Pembrolizumab	368	Advanced	33.7%, 32.9%	32.7	8.4
KEYNOTE-006	Ipilimumab	181	melanoma, ≤1 prior treatment	11.9%	15.9	3.4
	Nivolumab	272	Melanoma with	27%	16	3.1
CheckMate 037	Chemotherapy	herapy 133 progression on ipilimumab	10%	14	3.7	
T-VEC 295 Unresectable	26.4%	23.3	TTF: 8.2			
OPTIM	GM-CSF	141	melanoma	5.7%	18.9	TTF: 2.9

Robert, N Engl J Med 2015; Robert, Lancet 2019; Hodi, N Engl J Med 2010; Larkin, J Clin Oncol 2018.

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Trials in front-line melanoma

Trial	Treatment arm(s)	N	Patient selection criteria	ORR	Median PFS (months)	Landmark OS rate	Grade 3+ adverse events (%)
KEVNOTE 001	Dombrolizumah	GEE	Front-line	52%	16.9	5-year: 41%	170/
KETNOTE-001	Perindi Olizuinad	055	ITT	41%	8.3	5-year: 34%	1770
	Nivolumab + ipilimumab	314	Untreated stage III or IV	58%	11.5	5-year: 52%	59%
CheckMate 067	Nivolumab	316	melanoma	45%	6.9	5-year: 44%	23%
	Ipilimumab	315		19%	2.9	5-year: 26%	28%
	Nivolumab	210	Untreated BRAF WT	42.9%	5.1	3-year: 51.2%	15%
Checkiviate 066	Dacarbazine	208	advanced melanoma	14.4%	2.2	3-year: 21.6%	17.6%
IMspire150	Atezolizumab + cobimetinib + vemurafenib	256	BRAF V600 mutation- positive advanced/	66.3%	15.1	2-year: 60%	79%
	Cobimetinib + vemurafenib	258	metastatic melanoma	65.0%	10.6	2-year: 53%	73%

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- Consider combination ipilimumab/nivolumab up-front for patients with:
 - Brain metastases
 - Mucosal melanoma
 - High disease burden





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Question: How many combination doses to give



N=60	Week 6	Week 12	Best overall response rate
Overall response	35%	48%	57%
CR	0	5%	18%
PR	35%	43%	38%
SD	43%	18%	22%
PD	22%	30%	22%

Adverse events

- 100% of patients had any-grade irAEs, regardless of how many doses received
- 57% had grade 3-4 irAEs





Question: Does the sequence of targeted therapy and immunotherapy impact response?

Retrospective data suggests that patients who received BRAF inhibitors <u>prior</u> to treatment with pembrolizumab tended to have poorer outcomes on pembrolizumab therapy than those patients without prior BRAF inhibitor exposure.







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Adjuvant treatment options for melanoma

Drug	Indication	Dose
Dabrafenib + trametinib ⁺	Adjuvant BRAF+ melanoma with lymph node involvement following complete resection	Dabrafenib 150 mg twice daily + trametinib 2 mg daily
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
Ipilimumab*	Adjuvant therapy in stage III melanoma after complete resection	10 mg/kg Q3W for 4 doses, then 10 mg/kg Q12W for 3 years
Pembrolizumab	Adjuvant therapy of melanoma following complete resection – 1 year	200 mg Q3W or 400 mg Q6W
Nivolumab	Adjuvant treatment of melanoma after complete resection – 1 year	240 mg Q2W or 480 mg Q4W

⁺Not an immunotherapy; for reference

*not commonly used in this setting; historical reference



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Trials of adjuvant immunotherapy

Trial	Arms	Patient population	Ν	Key outcomes
FODTC 19071	Ipilimumab	Completely resected stage III	475	RFS HR: 0.76
EORIC 18071	Placebo	melanoma	476	OS HR: 0.72
EORTC 1325-	Pembrolizumab	High risk resected stage III	514	
MG/KEYNOTE-054	Placebo	melanoma	505	KF3 FIK. 0.30
CheckMate 238	Nivolumab	Resected stage IIIb or IV	453	
	Ipilimumab	melanoma	453	KF3 HK. 0.00
E1609	Ipilimumab 3 mg/kg		523	RFS HR: 0.85 OS HR: 0.78
	Ipilimumab 10 mg/kg	Resected stage IIIb-M1b melanoma	511	RFS HR: 0.84 OS HR: 0.88
	High-dose interferon alfa		636	





Adjuvant treatment considerations

- Goals of adjuvant treatment are different than goals of primary treatment
- Toxicity and quality of life are important considerations



Treatment-related adverse events





In development: Neoadjuvant immunotherapy in advanced melanoma

Trial	Regimen	N	pCR (%)	Median RFS (months)	Median follow-up (months)
Amaria Lancet Oncol 2018 (reference non-IO trial)	Dabrafenib + trametinib	21	58	19.7	18.6
Long Lancet Oncol 2019 (reference non-IO trial)	Dabrafenib + trametinib	35	49	23.0	27.0
Blank Nat Med 2018	lpilimumab + nivolumab	10	33	NR	32
	Nivolumab	12	25	NR	
Amaria Nat Med 2018	lpilimumab + nivolumab	11	45	NR	20
Huang Nat Med 2019	Pembrolizumab	30	19	NR	18
Rozeman Lancet Oncol 2019	lpilimumab + nivolumab	86	57	NR	8.3

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In development: Neoadjuvant immunotherapy in advanced melanoma

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Merkel cell carcinoma

- Associated with Merkel cell polyomavirus infection
- Higher incidence with weakened immune system (HIV, immunosuppressives) and increased age
- Distinct genomic profiles for UV- and virus-driven carcinomas
- Median PFS with chemo: ~90 days







Approved checkpoint inhibitors in Merkel cell carcinoma

Drug	Indication	Dose
Avelumab*	Patients >12 yr with metastatic Merkel cell carcinoma	800 mg Q2W + premedication (first 4 cycles)
Pembrolizumab	Adult/pediatric with recurrent advanced/metastatic Merkel cell carcinoma	Adults: 200 mg Q3W or 400 mg Q6W Pediatric: 2 mg/kg (up to 200 mg) Q3W

*Requires premedication with an antihistamine and acetaminophen prior to first four infusions





Avelumab in Merkel cell carcinoma

Setting	N	ORR	Median PFS	Median OS
First line	39	62.1%	9.1 months	
Second+ line	88	33.0%		12.6 months

First line



D'Angelo, JAMA Oncol 2018. D'Angelo, J Immunother Cancer 2020. © 2020–2021 Society for Immunotherapy of Cancer



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Second+ line



Pembrolizumab in 1st-line advanced Merkel cell carcinoma



Also an ongoing trial of adjuvant pembrolizumab for Merkel cell carcinoma (NCT03712605).









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Cutaneous squamous cell carcinoma

- Second-most common skin cancer
- Associated with high TMB and immunotherapy responsiveness







Approved checkpoint inhibitors for cutaneous squamous cell carcinoma

Drug	Indication	Dose
Cemiplimab-rwlc	Metastatic cutaneous squamous cell carcinoma, not candidate for curative therapies	350 mg Q3W
Pembrolizumab	Metastatic cutaneous squamous cell carcinoma	200 mg Q3W or 400 mg Q6W





Trials for R/M cutaneous SCC

Trial	Treatment	Ν	ORR	Median OS	Median PFS
KEYNOTE-629	Pembrolizumab	105	34.3%	NR	6.9 months
NCT02760498	Cemiplimab	59	47%	NR	NR





Pembrolizumab



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Approved checkpoint inhibitor for basal cell carcinoma

Drug	Indication	Dose
Cominlimah	Locally advanced BCC previously treated with hedgehog pathway inhibitor or for whom HHI is not appropriate	250 mg O 210/
Cemipiimab	Metastatic BCC previously treated with hedgehog pathway inhibitor or for whom HHI is not appropriate*	550 mg Q5 W
	*Accelerated approval	

Locally advanced disease	Metastatic disease
ORR: 29%	ORR: 21%
CR: 5/84	PR: 6/28
PR: 19/84	











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In development: Combination IO with BRAF targeted therapy



Multiple other triplet regimens are being tested.



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In development: Combination IO with oncolytic virus



Ribas et al Cell 2017



In development: Combination IO with pegylated 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



11 Molanoma (n-29 Efficacy Evoluable)	Overall Response	
IL Melanoma (n=38 Emcacy Evaluable)	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: Combination IO and TKI in mucosal melanoma

Treatment	N	ORR	Median PFS	Median OS
Toripalimab + axitinib	33	48.5%	7.5 months	20.7 months



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Strategies To Overcome Resistance



Incorporation of Biomarkers in Immunotherapy Clinical Trials





Conclusions

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





Additional Resources









Case Studies





Case Study 1

23yo F met melanoma of unknown primary to L groin. BRAF V600E+. PET showed NED elsewhere.

- s/p L groin LND and abd wall lesion excisional biopsy 10/16/2018. 5/20 LN +, abd wall lesion +.
- A neck nodule was biopsied 10/24/2018 and confirmed melanoma.
- Encorafenib + Binimetinib was started 11/7/2018. Serial scans showed stable neck nodules.
- Switched to nivolumab 480mg q4wks 9/23/2019.
- Resection of neck nodules 10/2/2019
- PET CT 11/18/2019 showed new R occipital brain met 1.4x1.1cm, R neck nodule 1.8x1cm and increased R inguinal nodes.

Question 1: What would you do, in addition to radiation therapy to the brain met?

- A. Enco/Bini
- B. Enco/Bini + anti-PD1
- C. Ipilimumab + nivolumab
- D. Hospice

Patient started ipi + nivo 11/21/2019, tolerated 2 doses. SRS to brain lets 12/9/2019. Treated stopped due to grade 3 hepatitis resolved with high dose steroids. PET CT and MRI brain 5/6/2020 showed generalized lymphadenopathy.

Question 2: What should be the next step?

- A. Enco/Bini
- B. Enco/Bini + anti-PD1
- C. lpi + nivo #3
- D. LN biopsy

Excisional biopsy of inguinal nodes 5/22/2020: benign nodes with non-caseating granulomas present. Maintainence nivo was started 6/2/2020. PET CT 8/3/2020 showed complete resolution of the lymphadenopathy. Patient's last dose nivo was on 1/19/2021. Last scan on the same day showed no evidence of active disease.





Case Study 2

60yo man noted one month of L eye dryness. Was found to have a eye lesion with rapid growth, now 1.5cm. Biopsy 8/2020 showed conjunctival melanoma, BRAF wild type. Whole body scan showed no disease elsewhere. Patient is seen by ophthalmology.

Question 1. What would you do?

- A. L orbit exenteration
- B. Resection of the lesion
- C. pembrolizumab
- D. Nivolumab + ipilimumab

Patient started pembrolizumab 400mg q6wks 9/2/2020, received 2 cycles. Had excision of the L conjunctival lesion on 9/4/2020 and repeated on 9/30/2020. Restaging CT on 11/13/2020 showed growth of the lesion to 2.4cm. No disease elsewhere.

Question 2 What is the next step?

- A. L orbit exenteration
- B. Resection of the lesion
- C. pembrolizumab
- D. Nivolumab + ipilimumab



11/9/2020



12/19/2020

1/26/2021

Nivo/Ipi was started on 11/19/2020, had three cycles so far and tolerating well. Patient is scheduled to have L orbital exenteration on 1/4/2021, but was found to have significant regression of the tumor.



Acknowledgements



THE HOPE FOUNDATION

Because answers to cancers come from clinical trials

SWOG Dr. Charles Coltman Award SWOG ITSC Pilot Award



5 For the Fight Fellow Award



Tower Cancer Research Foundation Career Development Award



ASCO Young Investigator Award ASCO Career Development Award



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SU2C/AACR Phil Sharp Innovation in Collaboration Award



UCLA CTSI KL2 Award



National Institutes of Health

SBIR R44 NCI Cancer Clinical Investigator Team Leadership Award (CCITLA)



MRA Young Investigator Award MRA Team Science Award

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