Immune checkpoint blockade

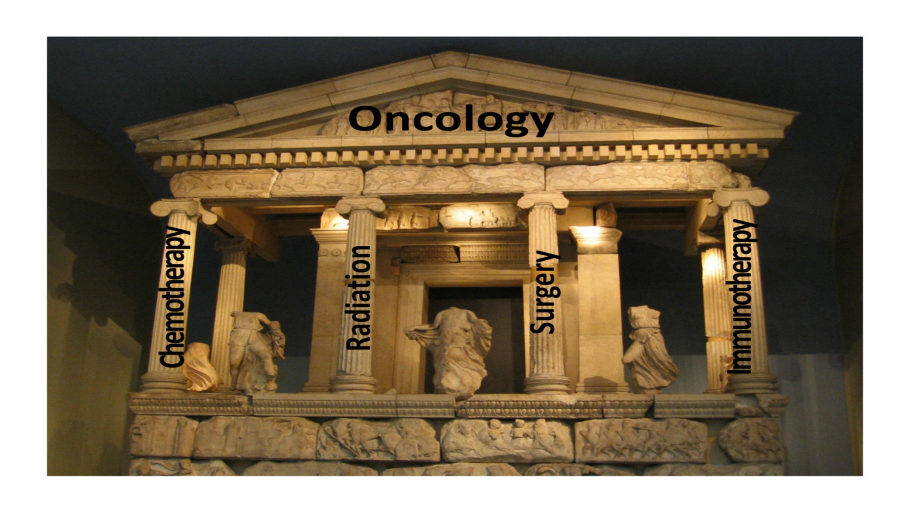
Immune Checkpoint Blockade

NCI CCR TRACO
Stephanie L. Goff, MD, FACS
September 19, 2022

Objectives

- The basics of immunotherapy
- Mechanism of action of checkpoint blockade
- Early clinical experience and the discovery of immune related adverse events
- Checkpoint blockade in melanoma
 - Ipilimumab
 - Nivolumab
 - Pembrolizumab
- Experimental Questions

Oncology

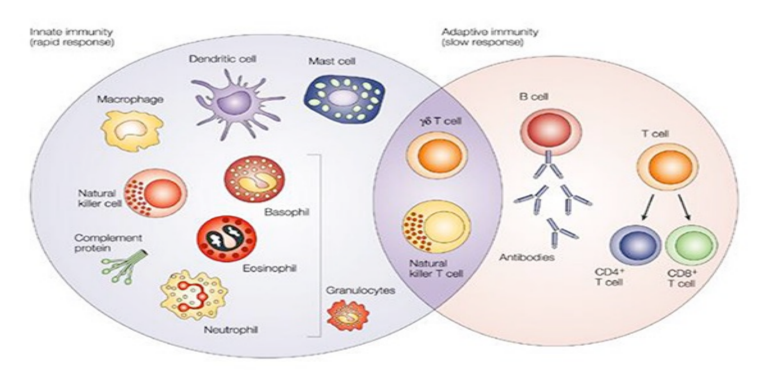


Cancer Immunotherapy

- 1. Nonspecific stimulation of immune reactions
 - a) Stimulate effector cells
 - b) Inhibit regulatory factors (checkpoint blockade)
- Active immunization to enhance anti-tumor reactions (cancer vaccines)
- 3. Passively transfer activated immune cells with antitumor activity (adoptive immunotherapy)

Immune system

Cells of the Immune System

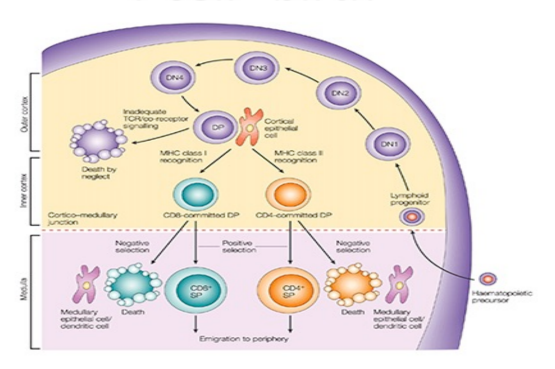


Nature Reviews | Cancer Dranoff 2004

Checkpoint blockade primarily affects T cells

T cell birth

T cell "birth"



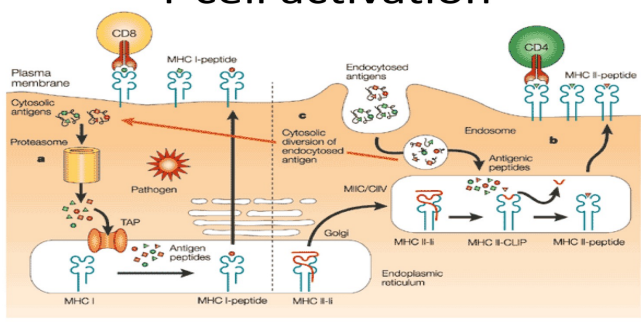
Nature Reviews | Immunology

Germain 2002

Builds a repertoire of T cells

T cell activation

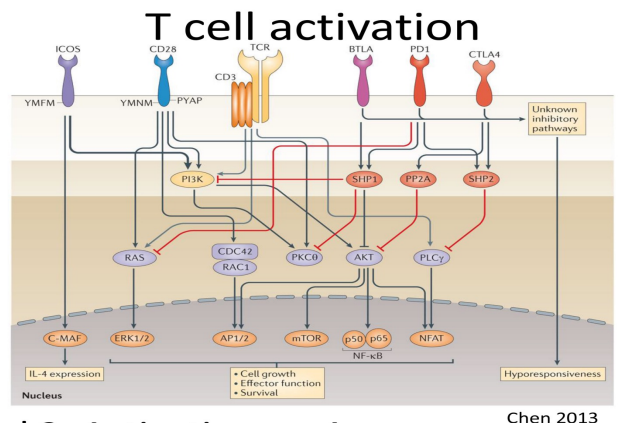
T cell activation



Nature Reviews | Immunology Heath 2001

- Signal 1: Specificity
- TCR engages antigen in context of MHC

T cell activation



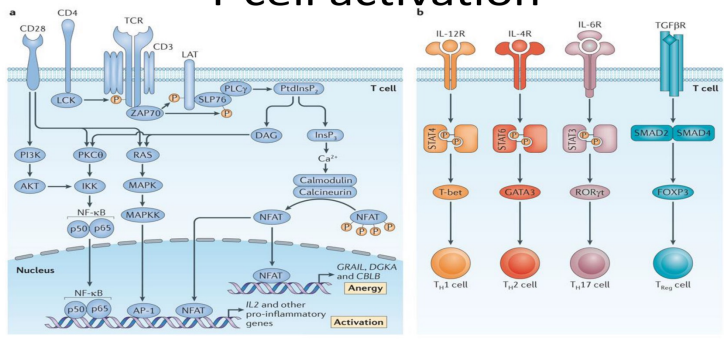
Signal 2: Activation vs. Anergy

Nature Reviews | Immunology

Costimulatory molecules

T cell activation

T cell activation



Nature Reviews | Immunology

Pollizzi 2014

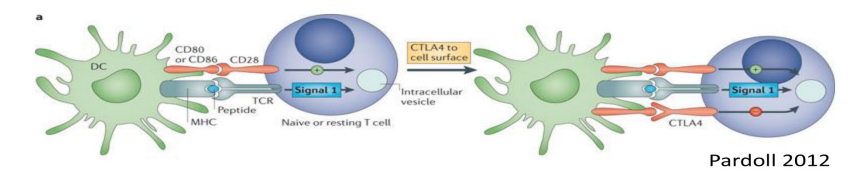
- Signal 3: Polarization
- Dependent on cytokine profile of the microenvironment

The role of Signal 2 checkpoints

- Immune checkpoints promote self-tolerance
 - Initial response to antigen occurs primarily in secondary lymphoid organs (lymph nodes, tonsils, spleen, Peyer's patches, mucosa associated lymphoid tissue)
- Immune checkpoints limit "collateral damage"
 - Effector recognition in peripheral tissue/tumor
- For cancer immunotherapy, two opportunities to break tolerance to self-antigen

CTLA-4

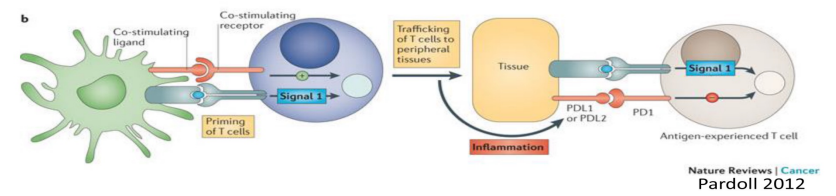
CTLA-4



- Naïve and memory T cells express surface CD28
- CTLA-4 is transported to the surface in correlation to the strength of CD28 stimulation
- CTLA-4 also competes with higher affinity for CD80/86
- A dampening effect on downstream processing
- Constitutively present on Treg cells

PD-1

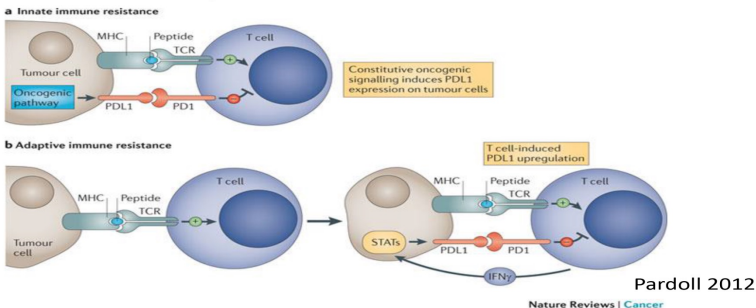
PD-1



- A primed T-cell is heading to peripheral tissue to engage a target, and once activated begin to express PD-1
- Inflammation present in the tissue can promote upregulation of the ligands of PD-1
- In general, this limits collateral damage during cellmediated destruction of infection

PD-1/PD-L1

PD-1/PD-L1 in cancer



- Cancer cells can increase the amount of PDL1
- Successful T-cell tumor destruction can increase PDL1 through upregulation in response to IFNγ

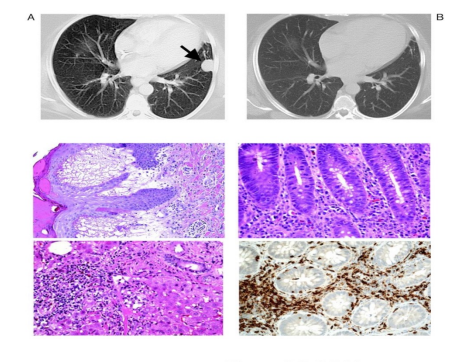
Checkpoint Blockade

- Where to start?
- Tumors known to respond to other immunotherapy
- Melanoma
- Estimated 9,940 deaths/year in US
- Metastatic disease
 16% 5 yr survival
- Interleukin-2 durable cure in 4%

- Renal Cell Cancer
- Estimated 14,080 deaths/year in US
- Metastatic disease
 12% 5 yr survival
- Interleukin-2 durable cure in 7%

Checkpoint Blockade @ NCI

- αCTLA-4, ipilimumab
- Phase I trial
- mAb (3mg/kg) + peptide
- Enrolled 14 patients
- 2 complete responders
- 1 partial response
- Accrual stopped for toxicity
 - Dermatitis, colitis, hepatitis, hypophysitis



Phan GQ 2003

PNAS

Checkpoint Blockade @ NCI

- Cautiously proceeded with Phase II trials in melanoma and RCC, initially with dose reduction (3 → 1 mg/kg)
- Objective response was associated with development of autoimmune events

Melanoma, p=0.008

	> Gr 3 AE	< Gr 3 AE
Objective Response (CR = 2)	5 (36%)	2 (5%)
Non-responder	9	40

RCC, p=0.009

	> Gr 3 AE	< Gr 3 AE
Objective Response (CR = 0)	5 (29%)	O (0%)
Non- responder	12	23

Attia P 2005

Yang JC 2007

Checkpoint Blockade @ NCI

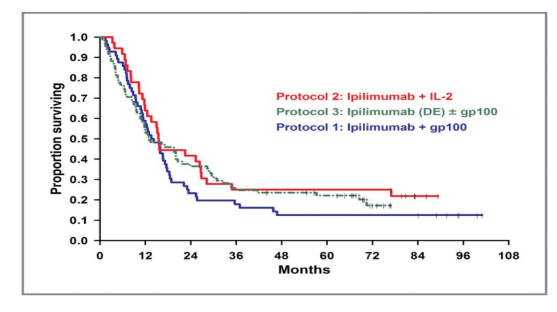
- Formal Phase II intrapatient dose escalation demonstrated association of response with immunerelated adverse events of any grade
- Enterocolitis was the most common grade 3/4 IRAE in patients with melanoma (18%) or RCC (28%)
- The administration of steroids to manage IRAE did not truncate responses

Melanoma, p=0.0004

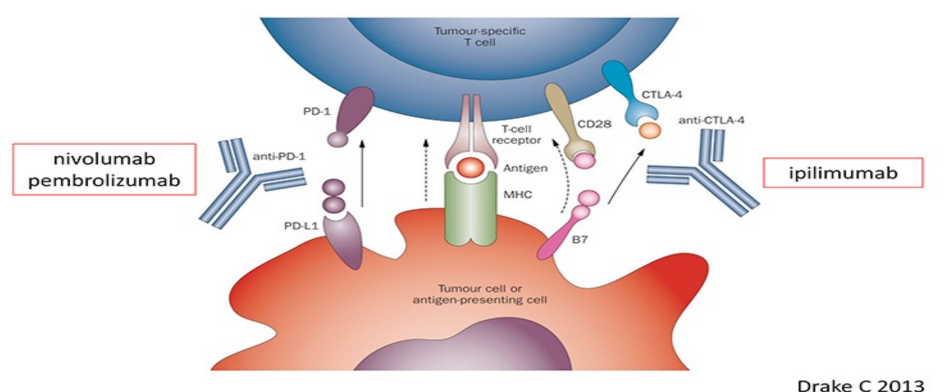
	Gr 3/4 IRAE	Gr 1/2 IRAE	No IRAE
Objective Response (CR = 3)	14 (28%)	8 (22%)	1 (2%)
Non- responder	36	28	52

Checkpoint Blockade @ NCI

- Developed algorithms for management of IRAEs
- Demonstrated durability of responses
 - OR 13-20%
 - 5 yr OS 13-23%



Checkpoint blockade in melanoma



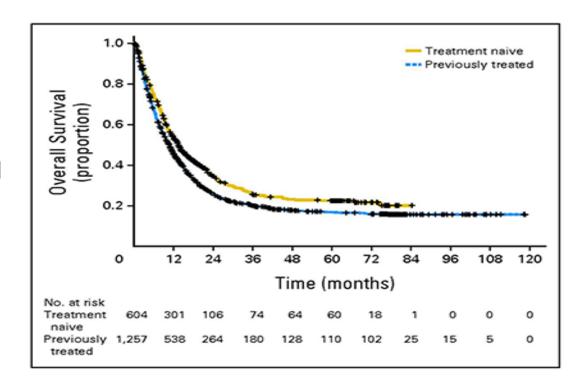
DIGIC CLINICAL

REVIEWS ONCOLOGY

Ipilimumab

Ipilimumab for melanoma

- Updated survival
- 3 year OS, 20-26%
- "Tail of the curve"
 - Durable for a small # of patients

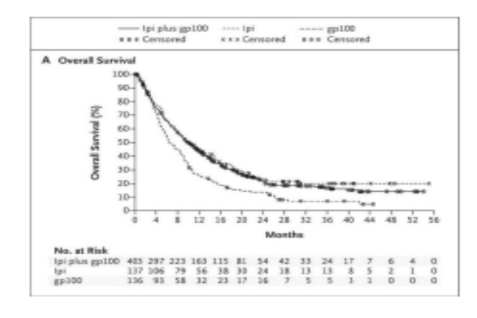


Ipilmumab

Ipilimumab for melanoma

- 11% response rate in Phase II trials at highest doses (10 mg/kg)
- Randomized Phase III
 ipilimumab ± gp100 vaccine vs.
 gp100 vaccine
- Allowed re-induction
- OR: ipilimumab arms
 7% (38/540)
 CR in 3 patients
- Disease control rate 22%
- Gr 3/4 irAE 10-15%

FDA approval for metastatic melanoma in March 2011



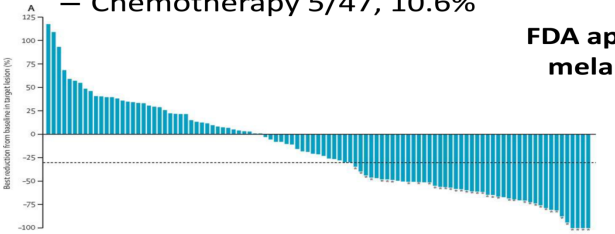


Nivolumab for melanoma

Nivolumab for melanoma

- Ipilimumab-refractory
- RCT: nivolumab vs chemotherapy of choice (CheckMate 037)
- Objective Response
 - Nivolumab 38/120, 31.7% with 4 CR





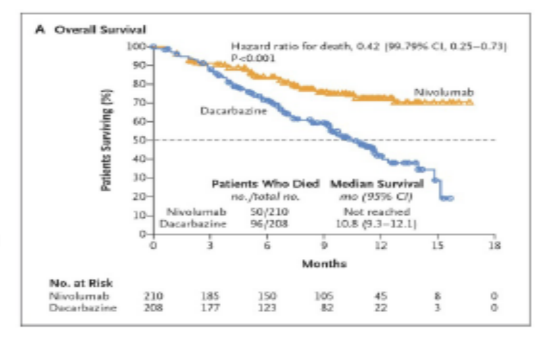
FDA approval for refractory melanoma in December 2014

> Weber IS 2015 THE LANCET Oncology

Nivolumab for melanoma

Nivolumab for melanoma

- Untreated metastatic disease
- Wildtype BRAF
- RCT: nivolumab vs dacarbazine (CheckMate 066)
- Objective response
 - Nivolumab 84/210 (40%)
 CR in 16 pts (7.6%)
 - Dacarbazine 29/208 (14%) CR in 2 pts (1%)



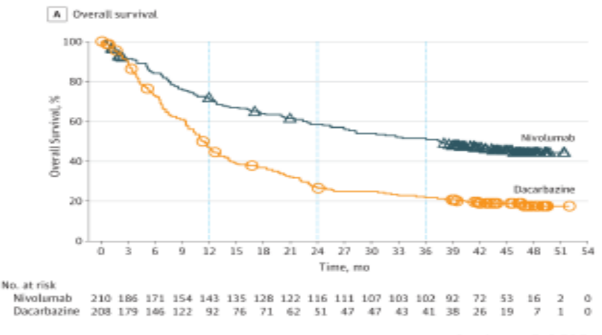
Approved for initial treatment (BRAF-wt) in November 2015



Nivolumab

Nivolumab for melanoma

- Overall Survival update for Checkmate 066
- Three-year OS:
 - Nivolumab 51%
 - Dacarbazine 22%

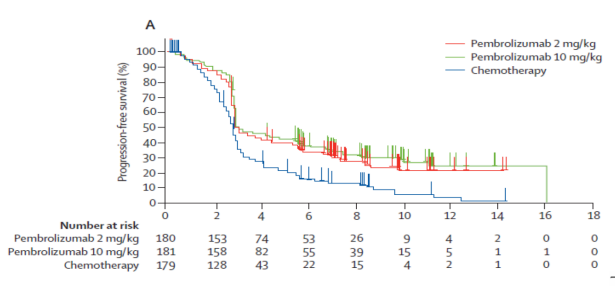


Ascierto P 2018

Pembrolizumab for melanoma

Pembrolizumab for melanoma

- Ipilimumab-refractory
- Phase II, dose comparison (2mg/kg vs 10 mg/kg) vs chemo
- 540 patients
 - 2mg/kg ORR 38 (21%), 10 mg/kg ORR 46 (25%), chemo 8 (4%)
- Grade 3/4 AE 12%



Weber JS 2015

Pembrolizumab for melanoma

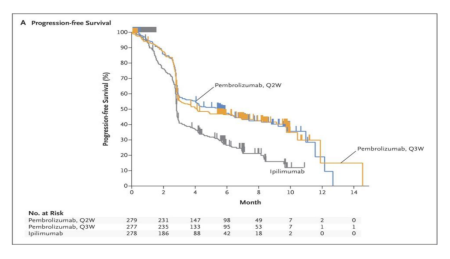
Pembrolizumab for melanoma

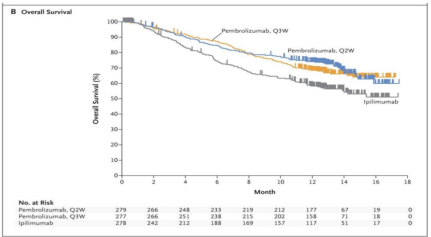
- RCT, KEYNOTE-006, first-line therapy
- Pembrolizumab (q2w, q3w) vs ipilimumab
- 1:1:1
- 834 patients
- Objective Response
 - Pembrolizumab q2w 94/279 (33.7%), CR 14
 - Pembrolizumab q3w 91/277 (32.9%), CR 17
 - Ipilimumab 33/278 (11.9%), CR 4



Pembrolizumab for melanoma

Pembrolizumab for melanoma





Grade ≥3 AE

- Pembrolizumab q2w 13.3% (1.4% Colitis)
- Pembrolizumab q3w 10.1% (2.5% Colitis)
- Ipilimumab 19.9% (7% Colitis)

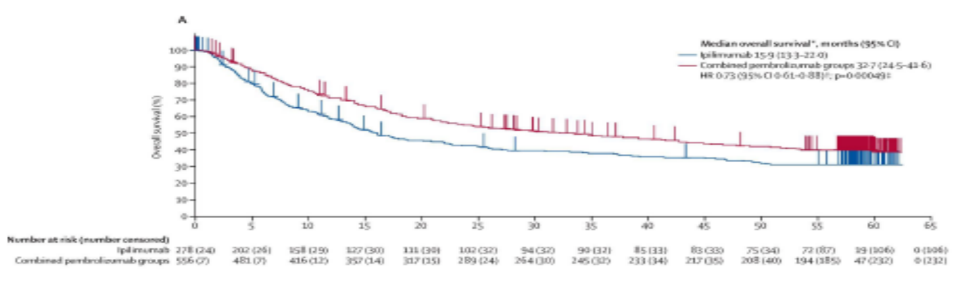
Robert C 2015

The NEW ENGLAND
JOURNAL of MEDICINE

Pembrolizumab

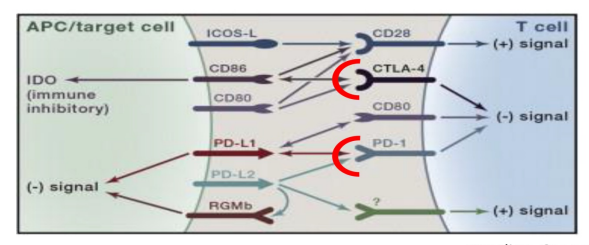
Pembrolizumab for melanoma

Three year OS of 48.1% vs 37.8%



Checkpoint modulation

Checkpoint Modulation



Topalian, Cancer Cell 2015

- In melanoma, the two approved antibodies interfere with separate receptor/ligand complexes
- Could combination therapy improve response or survival?

Nivolumab/Ipilmumab

Nivolumab/Ipilimumab for melanoma

- Previously untreated
- Phase III, RCT
- 945 patients
- 1:1:1
- PD-L1 (+) ≥5%

Characteristic	Nivolumab (N=336)	Nivolumab plus lpilimumab (N=314)	lpilimumab (N=315)	Total (N = 945)
PD-L1 status — no. (%)				
Positive	80 (25.3)	68 (21.7)	75 (23.8)	223 (23.6)
Negative	208 (65.8)	210 (66.9)	202 (64.1)	620 (65.6)
Could not be determined or evaluated	28 (8.9)	36 (11.5)	38 (12.1)	102 (10.8)
BRAF status — no. (%)				
Mutation	100 (31.6)	101 (32.2)	97 (30.8)	298 (31.5)
No mutation	216 (68.4)	213 (67.8)	238 (69.2)	647 (68.5)

Nivolumab/Ipilmumab

Nivolumab/Ipilimumab for melanoma

- Previously untreated
- Phase III, RCT
- 945 patients
- 1:1:1
- Grade 3/4 AE
 - Nivolumab 16.3%
 - Ipilimumab 27.3%
 - Combo 55.0%

/ariable	Nivolumab (N=316)	Nivolumab plus Ipilimumab (N = 314)	Ipilimumab (N=315)
Best overall response — no. (%)*			
Complete response	28 (8.9)	36 (11.5)	7 (2.2)
Partial response	110 (34.8)	145 (46.2)	53 (16.8)
Stable disease	34 (10.8)	41 (13.1)	69 (21.9)
Progressive disease	119 (37.7)	71 (22.6)	154 (48.9)
Could not be determined	25 (7.9)	21 (6.7)	32 (10.2)
Objective response†			
No. of patients with response	138	181	60
% of patients (95% CI)	43.7 (38.1–49.3)	57.6 (52.0-63.2)	19.0 (14.9–23.8)
Estimated odds ratio (95% Ci);	3.40 (2.02–3.72)	6.11 (3.59–10.38)	
Two-sided P value	< 0.001	< 0.001	2 —
Time to objective response — mo			
Median	2.78	2.76	2.79
Range	2.3-12.5	1.1-11.6	2.5-12.4

^{*} The best overall response was assessed by the investigator according to the Response Evaluation Criteria in Solid Tumors, version 1.1.

Larkin J 2015



[†] Data included patients with a complete response and those with a partial response. The calculation of the confidence interval was based on the Clopper–Pearson method. These analyses were conducted with the use of a two-sided Cochran–Mantel–Haenszel test stratified according to PD-L1 status, BRAF mutation status, and metastasis stage.

[‡] The comparison is with the ipilimumab group.

Nivolumab/Ipilmumab

Nivolumab/Ipilimumab for melanoma

- Previously untreated
- Phase III, RCT
- 945 patients
- 1:1:1
- Grade 3/4 AE
 - Nivolumab 21%
 - Ipilimumab 28%
 - Combo 59%

Variable	Nivolumab plus Ipilimumab (N = 314)	Nivolumab (N=316)	(N = 315)
Best overall response — no. (%)†			
Complete response	61 (19)	52 (16)	16 (5)
Partial response	122 (39)	88 (28)	43 (14)
Stable disease	38 (12)	31 (10)	69 (22)
Progressive disease	74 (24)	121 (18)	159 (50)
Unable to determine	19 (6)	24 (8)	28 (9)
Objective response:			
No. of patients with response	183	140	3.9
% of patients (95% CI)	58 (53-64)	44 (39-50)	19 (15-24)
Estimated adds natio (95% CI)§	6.46 (4.45-9.38)	3.57 (2.48-5.15)	-
Pivalue	<0.001	-0.001	
Median duration of response (95% CI) - mo	NR	NR (36.3-NR)	19.3 (8.3-NR)

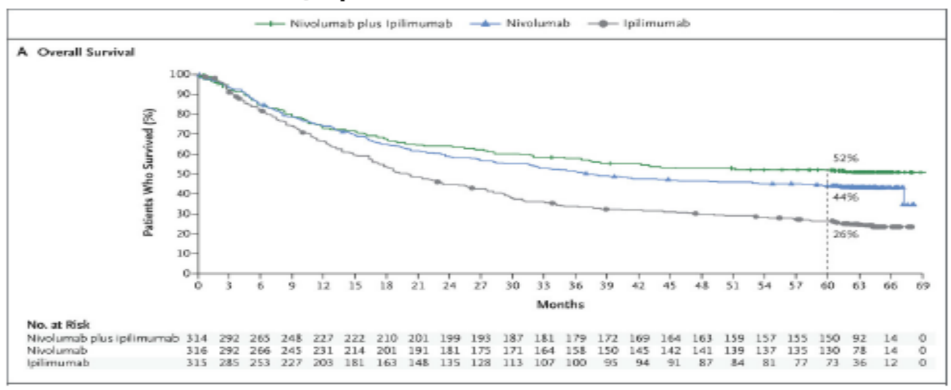
FDA approval of combination for melanoma in January 2016

Wolchok J 2017



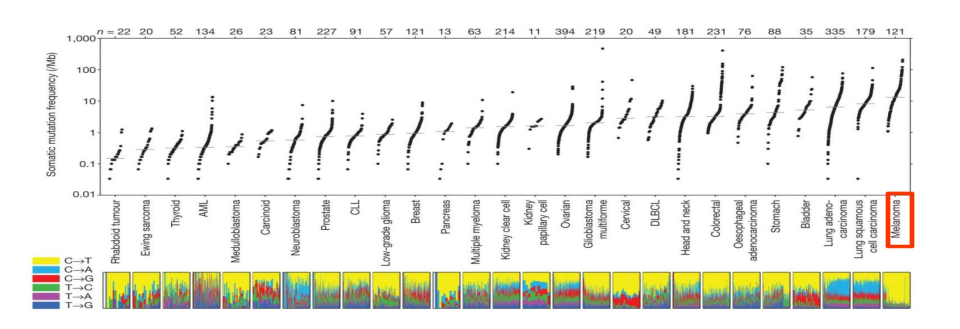
Nivolumab/Ipilmumab for melanoma

Nivolumab/Ipilimumab for melanoma



Melanoma

Why melanoma?



Highly mutated tumors

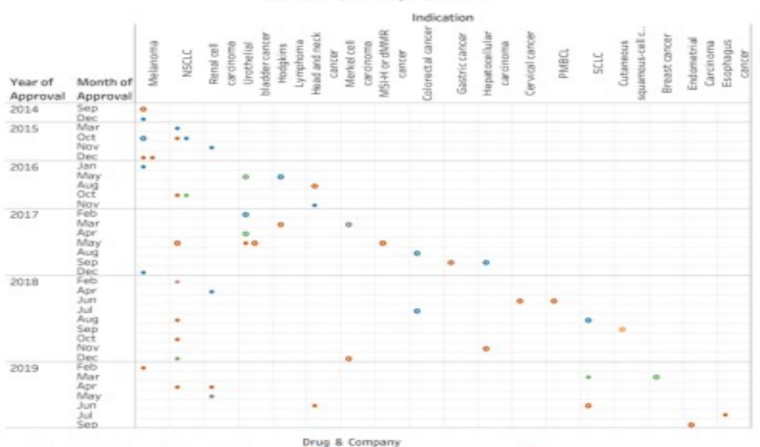
- Non-small cell lung cancer
- ~158,040 deaths/year in US
- Regional disease
 16% 5 yr survival
- Metastatic disease2% 5 yr survival
- Correlation between smoking and # mutations

- Tumors with mismatch repair (MMR) deficiency
 - Lynch syndrome (germline mutation)
 - Sporadic mutation
 - MSH2, MLH1, MSH6, PMS2
- Bladder cancer
 - 16,000 deaths/year in US
 - Highly lethal once metastatic

Timeline

Timeline of Anti-PD-1/L1 Antibody Approvals by the FDA

Updated on September 23, 2019, by Jun Tang/Annie Yu Sources: CRI, CRI Analytics, and FDA



Pembrolizumab, Merck Co.

Accelerator Accelerator Accelerator

Nivolumab, Bristol-Myers Squibb

Durvalumab, AstraZeneca

Cemiplimab, Regeneron

Avelumab, Pfizer/Merck KGaA

The Anna-Maria Kellen

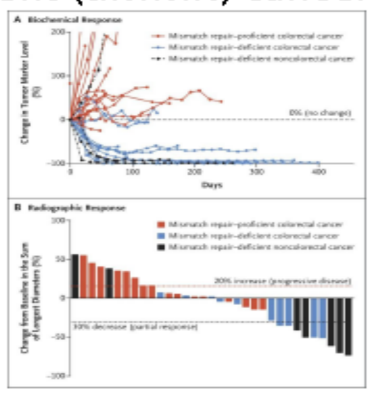
Clinical

CANCER

Mismatch/repair deficiency

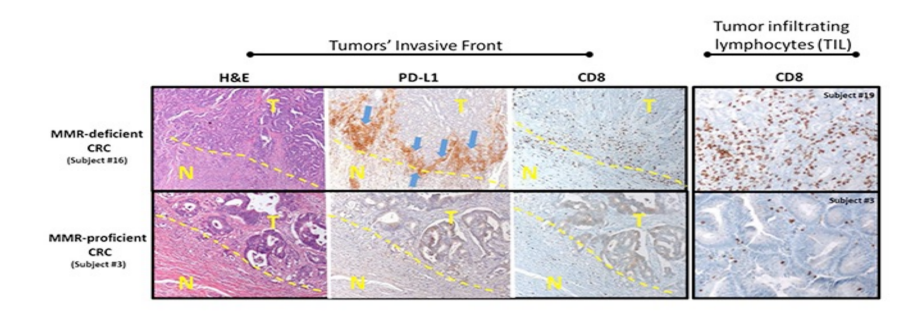
Pembrolizumab for mismatch repair deficient (dMMR) cancer

- Builds on hypothesis of neoantigens from somatic mutations
- Phase 2 study
- Three parallel cohorts
 - MMR-proficient CRC
 - MMR-deficient CRC
 - MMR-deficient other



Tumor-stromal interface

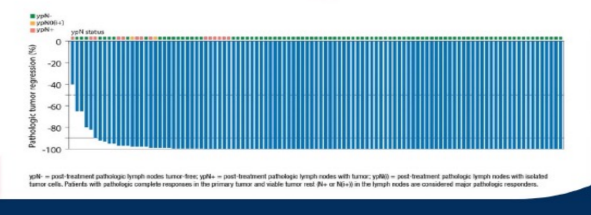
Pembrolizumab at the tumor-stroma interface



Pre-op combinations checkpoint

Pre-op combination checkpoint





Neoadjuvant immunotherapy in dMMR colon cancer - a paradigm shift?



Checkpoint blockade

Checkpoint Blockade

- Highly mutated tumors
 - Melanoma
 - Non-small cell lung cancer
 - Bladder cancer
 - Tumors with mismatch repair deficiency

- Use in other tumors?
 - Renal cell
 - Responds to other immun otherapy
 - Hodgkin's lymphoma
 - Reed-Sternberg cells have elevated amounts of PD-L1
 - Head and neck SCC
 - HPV and mutations

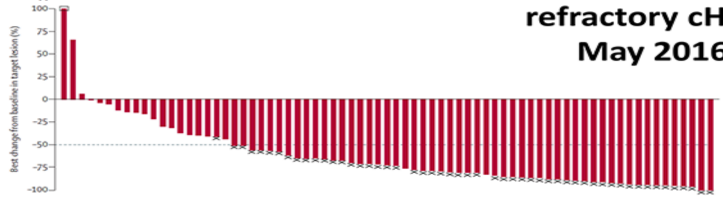
Hodgkin's lymphoma

Nivolumab for Hodgkin's Lymphoma

- 80 patients
 - Refractory to stem cell transplant
 - Refractory to brentuximab

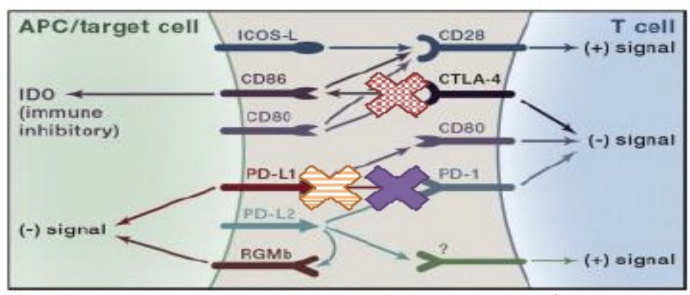
- Objective Response
 - 53/80 (66%)
 - 7 complete remission

FDA approval for refractory cHL in May 2016



Checkpoint modulation

Checkpoint Modulation



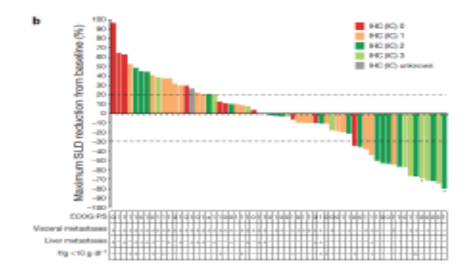
To pe lien, Cance r Cell 2015

- Initial focus on blocking Signal 2 on the T cell side
 - Anti-CTLA-4: ipilimum ab (Yervoy), tremelimum ab
 - 💓 Anti-PD-1: nivolumab (Opdivo), pembrolizumab (Keytruda), cemiplimab (Libtayo).
- Newer development on blocking Signal 2 on the target
 - Anti-PD-L1: atezolizumab (Tecentriq), avelumab (Bavencio), durvalumab (Imfinzi)

Bladder cancer

αPD-L1 in Urothelial bladder cancer

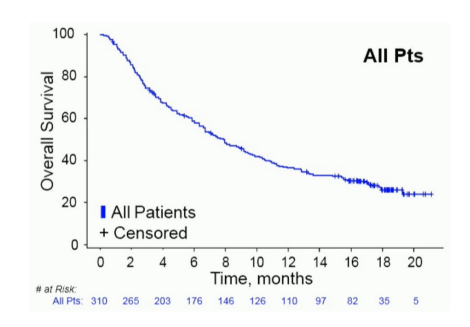
- MPDL3280A
- Atezolizumab
- 15 mg/kg q3w
- 27% tumors with >5% PD-L1 by IHC
- 65 patients with pretreatment biopsy
- Objective Response
 - ≥ 5% PD-L1 13/30 (43.3%)
 - < 5% PD-L1 4/35 (11.4%)</p>
- Grade 3/4 AE 4%





αPD-L1 in Urothelial bladder cancer

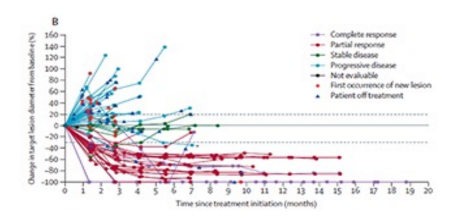
- 310 patients
- Objective Response
 - -45 (15%)
 - With 15 complete responses
- Overall Survival
 - -7.9 months
- 1 yr Survival
 - -37%



Avelumab

Avelumab in Merkel cell carcinoma

- 88 patients
 - Confirmed metastatic disease
- Objective Response
 - **28/88 (32%)**
 - 8 complete remission



FDA approval for Merkel cell carcinoma in March 2017

PD-1/PD-L1 pathway

Blocking the PD-1/PD-L1 pathway

	Drug	Melanoma	NSCLC	RCC	Bladder
Anti-PD-1	Nivolumab	32% (n=107)	17% (n=129) 30% (n=20)	29% (n=34) 21% (n=168)	NR
	Pembrolizumab	38% (n=135) 26% (n=157)	26% (n=42) 20% (n=194)	NR	24% (n=29)
Anti-PD-L1	BMS-936559	17% (n=52)	10% (n=49)	12% (n=17)	NR
	MEDI4736	NR	16% (n=58)	NR	NR
	Atezolizumab	30% (n=43)	23% (n=53)	14% (n=56)	26% (n=65)

FDA Approved (As of 9/2016)

Adapted from Lipson 2015



Altezolizumab

Atezolizumab (αPD-L1) for melanoma

- BRAF V600E/K mutation
- Phase III RCT, with BRAK/MEK inhibitors
- 514 patients, randomized 1:1

cobimetinib cobimetinib Progression-free survival, months. 15 1 (11-4-18-4) 10-5 (9-3-12-7) median (95% O) 90 FDA approval of Hazard ratio (95% CI) 0.78 (0.63-0.97), log-rank p=0.0249 80-(X) powers out-mossessor 70combination for 60-50melanoma in 40 -45.1% 30 - Placebo » vernusafenib » cobirnetinib 20-July 2020 Atezokzumab + vemurafenib + cobimetinib 10-Censored 12 á 24 Time (months) Murebor attrisk Placebo +vernurafenib +cohimetini b 230 143 987 Atecolizamab + vernurafenib + cobimetini b 255 229 174 149 123

Gutzmer R 2020 THE LANCET

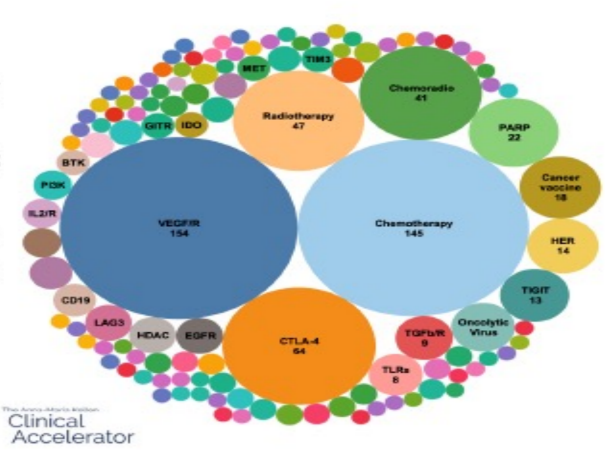
Ategoligumab + vemurafenib +

Placebo + verrorafenib +

Combination clinical trials

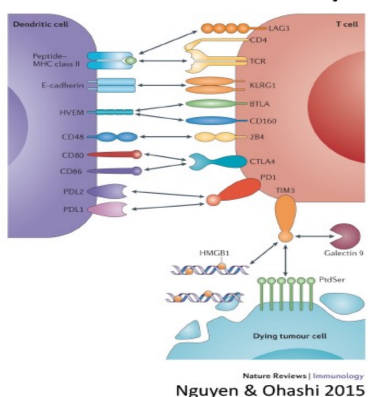
Combination Clinical Trials

- Over 2900
 different trials of
 combination
 therapy with 253
 different agents
- 724 new trials in first 9 months of 2020



New checkpoint inhibitors

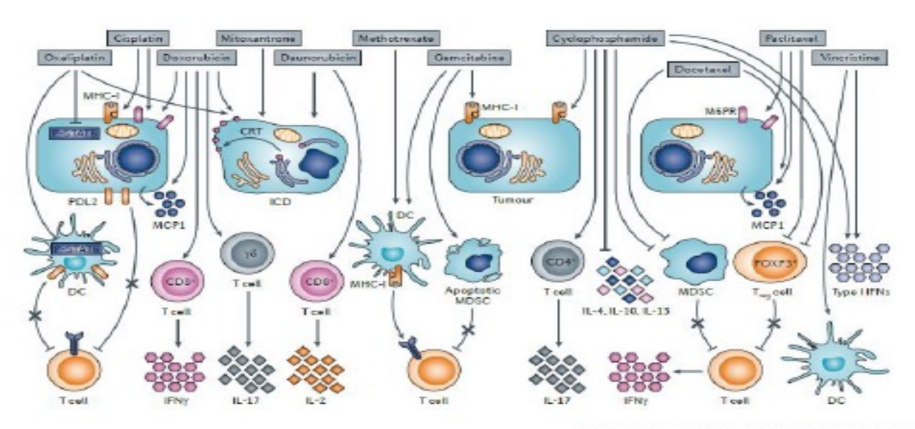
New checkpoint inhibitors



- LAG-3
- Combination formula
 - Anti PD-1
 - Anti LAG-3
- 16% complete response
- 27% partial respons
- Approved for 1st line metastatic melanoma in March 2022

Chemotherapy combinations

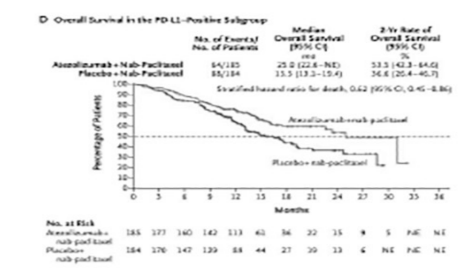
Rationale for Chemotherapy Combinations



Breast cancer

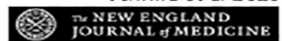
PD-L1/chemo in mBrCa

- Nab-paclitaxel ± atezolizumab
- 902 patients
- Randomized
- 379 with PD-L1+ (≥1%) tumors
- Objective Response
 - Chem o + atezo 59%
 - Chem o + placebo 43%
- 2yr Survival
 - Chemo + atezo 54%
 - Chem o + placebo 37%



FDA approval for PD-L1+ TNBC in March 2019

Schmid et al 2018



Checkpoint modulators

Checkpoint Modulators

 Every expanding list of indications

 Over 2200 different trials of combination therapy

Any questions?

