

Immunotherapy for the Treatment of Head and Neck Cancer

Greg Durm, MD

Assistant Professor of Clinical Medicine Indiana University Simon Cancer Center









Disclosures

- Merck- Research Funding
- Bristol-Myers Squibb- Research Funding
- Astra Zeneca- Research Funding

• I will not be discussing non-FDA approved indications during my presentation.



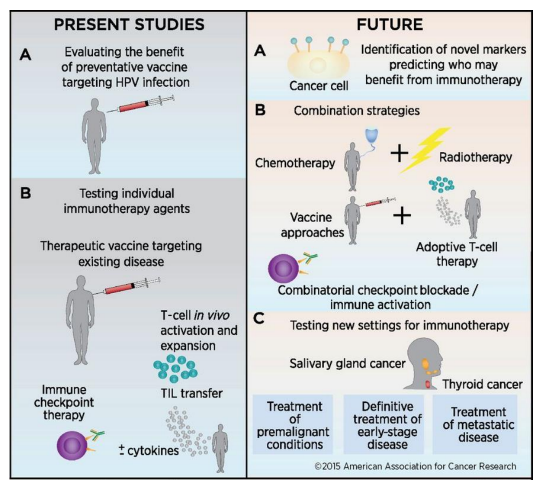


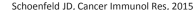




Immunotherapy for the Treatment of Head and Neck Cancers

- Immuno-Oncology (I-O) developments in treatment of head and neck cancers
 - Expression of immunologic markers to guide treatment
 - Preventive vaccination against virally mediated cancers
 - PD-1 checkpoint inhibitors for the treatment of metastatic disease







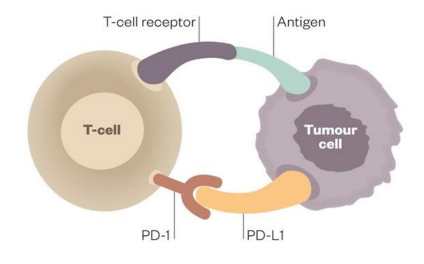




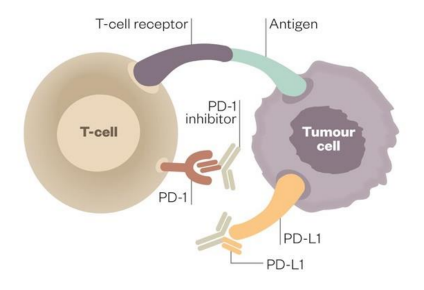


Immunotherapy for the Treatment of Head and Neck Cancers

Immune Checkpoint Inhibitors (ICI)



PD-1 acts as "off-switch" for T cells, allowing cancer cells to evade immune attack



Antibodies against PD-1 and PD-L1 boost the immune response against cancer cells









FDA-approved Checkpoint Inhibitors for Use in Head and Neck Cancers

- Pembrolizumab 200 mg IV Q3W(anti-PD-1)
 - KEYNOTE 012/055: Patients with recurrent or metastatic (R/M) squamous cell carcinoma of the head and neck (HNSCC) with disease progression on or after platinum-containing chemotherapy
 - Accelerated Approval by FDA August 5, 2016
- Nivolumab 240 mg IV Q2W or 480 mg IV Q4W (anti-PD-1)
 - CheckMate 141: Patients with R/M HNSCC with disease progression on or after a platinumbased therapy
 - Breakthrough Therapy Designation by FDA April, 2016
 - Approval November 10, 2016







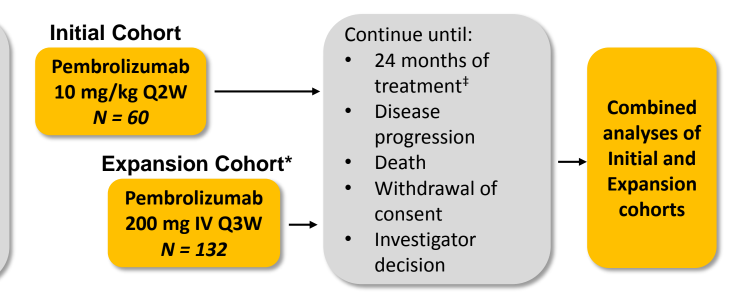


KEYNOTE-012: Pembrolizumab in R/M HNSCC

Nonrandomized, Phase 1b Trial, Cohorts[†] B, B2

Patients

- R/M HNSCC
- Measurable disease (RECIST v1.1)
- ECOG PS 0-1
- PD-L1+ (initial cohort)
- PD-L1+ or PD-L1-(expansion cohort)



Response assessment: Every 8 weeks until disease progression

Primary end points: ORR (RECIST v1.1, central imaging vendor review), safety

Secondary end points: ORR (investigator), PFS, OS, duration of response (DOR), ORR in HPV+ patients§







[†]Additional cohorts included bladder cancer, TN breast cancer, and gastric cancer.

[‡]Treatment beyond progression was allowed.

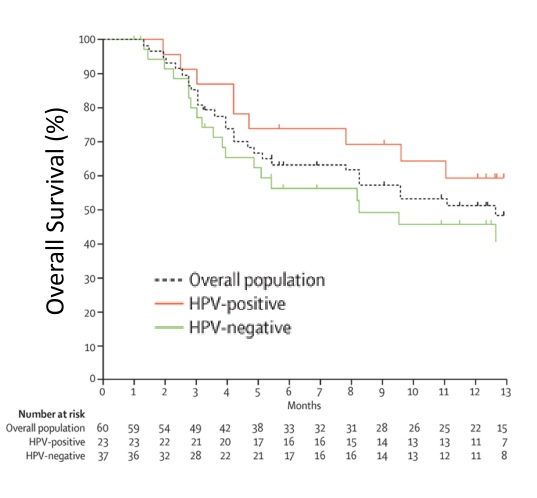
[§] Initial cohort only.

^{*}Median duration of disease not reached.



KEYNOTE-012: Pembrolizumab in R/M HNSCC

Nonrandomized, Phase 1b Trial, Cohorts[†] B, B2



- ORR = 18%
 - CR = 4%
 - PR = 14%
- mOS = 8.0 months
- mPFS = 2.2 months

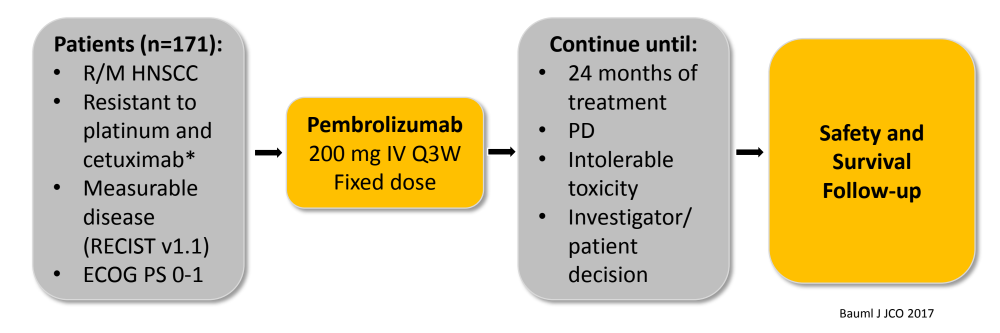








KEYNOTE-055: Pembrolizumab in R/M HNSCC after Progression on Platinum/Cetuximab Phase II Trial, Single Arm



Response assessment: Imaging every 6 to 9 weeks (central radiology review)

Primary end points: ORR (RECIST v1.1) by Response Evaluation Criteria in Solid Tumors and safety

Secondary end points: ORR (RECIST v1.1) in all dosed patients, ORR for HPV+, PD-L1+, DOR, PFS, OS

*75% of patients had ≥ 2 prior lines of therapy for metastatic disease









KEYNOTE-055: Pembrolizumab in R/M HNSCC after Progression on Platinum/Cetuximab Phase II Trial, Single Arm

Outcome	All Patients N=171	HPV Status		PD-L1 Status		
		Positive n=37	Negative n=131	≥1% n=140	<1% n=26	≥50% n=48
ORR, %	16	16	15	18	12	27
mPFS, mo	2.1					
6-mo PFS,%	23	25	21	24	20	31
6-mo OS, %	59	72	55	59	56	60

- Neither tumor PD-L1 expression or HPV status are sufficiently robust in guiding the use of pembrolizumab at this time.

Bauml J, et al, J Clin Oncol. 2017









CheckMate 141: Nivolumab vs Investigator's Choice in R/M HNSCC after Platinum Therapy Phase III Randomized, Safety and Efficacy Trial

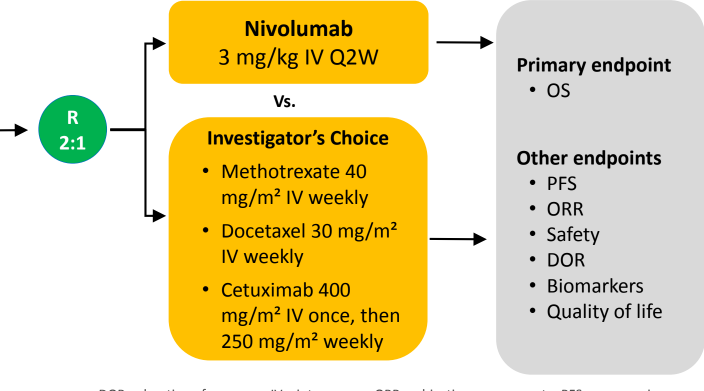
Key Eligibility Criteria

- R/M SCCHN of the oral cavity, pharynx, or larynx
- Progression on or within 6 months of last dose of platinum-based therapy
- Irrespective of no. of prior lines of therapy
- Documentation of p16 to determine HPV status (oropharyngeal)
- Regardless of PD-L1 status^a

Stratification factor

Prior cetuximab treatment

^aTissue required for testing



DOR = duration of response; IV = intravenous; ORR = objective response rate; PFS = progression-free survival; Q2W = once every 2 weeks; R = randomized. Clinicaltrials.gov NCT02105636.

Ferris & Gillison, NEJM, 2016



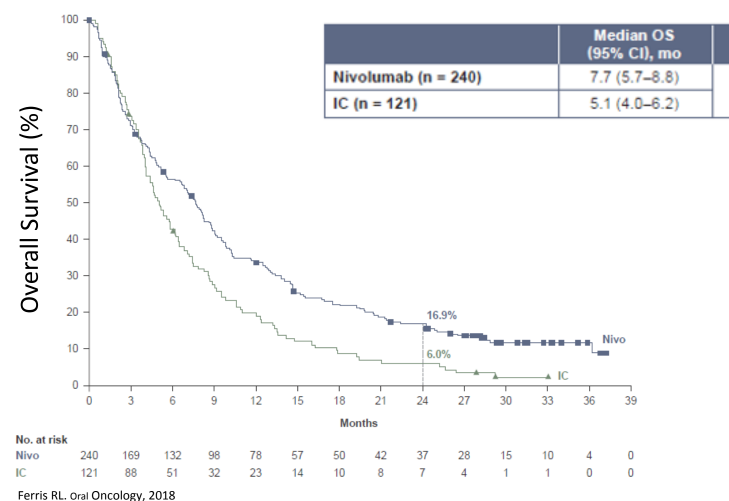






Checkmate 141: Nivolumab vs Investigator's Choice in R/M HNSCC after Platinum Therapy

Overall Survival: 2 year report







HR

(95% CI)

0.68

(0.54 - 0.86)

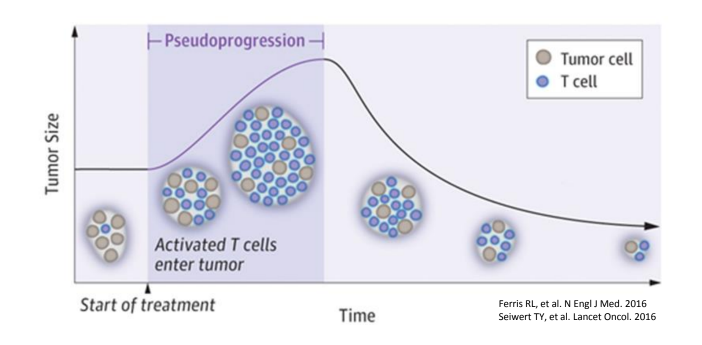




Response to Immune Checkpoint Inhibitor Treatment with Brief Increase in Tumor Size

Pseudoprogression

- Early appearance of an increase in tumor burden, subsequently followed by tumor regression
- Initially recognized in the melanoma trials, with incidence up to 10%











Response to Immune Checkpoint Inhibitor Treatment with Brief Increase in Tumor Size

Case Report – KEYNOTE-012



 Both KEYNOTE-012 and CheckMate 141 trials showed an exceedingly rare rate of pseudoprogression with pembrolizumab and nivolumab, respectively.

> Ferris RL, et al. N Engl J Med. 2016 Seiwert TY, et al. Lancet Oncol. 2016









Evaluating Biomarkers in HNSCC

- Current FDA approvals of pembrolizumab and nivolumab are NOT contingent upon tumor PD-L1 status
 - KEYNOTE 012/055: Response rates not significantly different on the basis of tumor PD-L1 staining
 - KEYNOTE 040: Phase III pembrolizumab vs. investigator's choice chemotherapy
 - Did not meet survival endpoints in total population but improved outcomes in patients with PD-L1 expressing tumor
 - CheckMate 141: Most benefit was seen in PD-L1-positive tumors



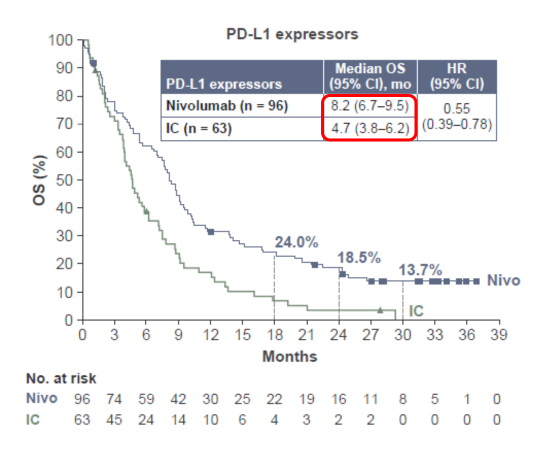


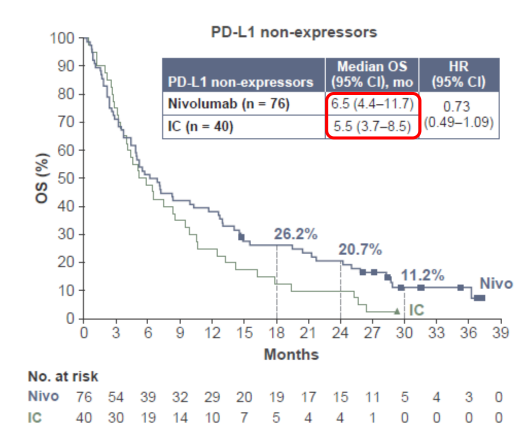




Evaluating Biomarkers in HNSCC

CheckMate 141: 2 year update













Immune-related Adverse Events

KEYNOTE 012

Table 2. Treatment-Related Adverse Events by Grade Severity (all-patients-astreated population: N = 132)

treated population; N = 132)				
Treatment-Related Adverse Event	Grade 1 or 2 (≥ 10% of patients), No. (%)	Grade 3 (any occurrence), No. (%)	Grade 4 (any occurrence), No. (%)	
Patients with ≥ 1 event	70 (53)	8 (6)	4 (3)	
Hypothyroidism	14 (11)	0	0	
Immune thrombocytopenic purpura	0	0	1 (1)	
Abdominal pain	1 (1)	1 (1)	0	
Colitis	0	1 (1)	0	
Dysphagia	1 (1)	1 (1)	0	
Nausea	6 (5)	1 (1)	0	
Stomatitis	1 (1)	1 (1)	0	
Facial edema	0	1 (1)	0	
Fatigue	28 (21)	0	0	
Localized edema	0	1 (1)	0	
Infection	0	1 (1)	0	
Decreased appetite	9 (7)	2 (2)	0	
Dehydration	0	1 (1)	0	
Diabetic ketoacidosis	0	0	1 (1)	
Hyperglycemia	1 (1)	0	1 (1)	
Type I diabetes mellitus	0	1 (1)	0	
Laryngeal edema	0	0	1 (1)	
Pneumonitis	2 (2)	2 (2)	0	
Respiratory distress	0	1 (1)	0	
Facial swelling	3 (2)	1 (1)	1 (1)	
_				

CheckMate 141

Event	Nivolumab (N=236)		
	Any Grade	Grade 3 or 4	
Any event	139 (58.9)*	31 (13.1)	
Fatigue	33 (14.0)	5 (2.1)	
Nausea	20 (8.5)	0	
Rash	18 (7.6)	0	
Decreased appetite	17 (7.2)	0	
Pruritus	17 (7.2)	0	
Diarrhea	16 (6.8)	0	
Anemia	12 (5.1)	3 (1.3)	
Asthenia	10 (4.2)	1 (0.4)	
Vomiting	8 (3.4)	0	
Dry skin	7 (3.0)	0	
Stomatitis	5 (2.1)	1 (0.4)	
Weight loss	4 (1.7)	0	
Mucosal inflammation	3 (1.3)	0	
Peripheral neuropathy	1 (0.4)	0	
Alopecia	0	0	
Neutropenia	0	0	









Immune-related Adverse Events

Table 2 General guidance for corticosteroid management of immune-related adverse events

Grade of immune-related AE (CTCAE/equivalent)	Corticosteroid management	Additional notes
1	Corticosteroids not usually indicated	Continue immunotherapy
2	 If indicated, start oral prednisone 0.5-1 mg/kg/day if patient can take oral medication. If IV required, start methylprednisolone 0.5-1 mg/kg/day IV If no improvement in 2–3 days, increase corticosteroid dose to 2 mg/kg/day Once improved to ≤grade 1 AE, start 4–6 week steroid taper 	 Hold immunotherapy during corticosteroid use Continue immunotherapy once resolved to ≤grade 1 and off corticosteroids Start proton pump inhibitor for GI prophylaxis
3	 Start prednisone 1-2 mg/kg/day (or equivalent dose of methylprednisolone) If no improvement in 2–3 days, add additional/alternative immune suppressant Once improved to ≤ grade 1, start 4–6-week steroid taper Provide supportive treatment as needed 	 Hold immunotherapy; if symptoms do not improve in 4–6 weeks, discontinue immunotherapy Consider intravenous corticosteroids Start proton pump inhibitor for GI prophylaxis Add PCP prophylaxis if more than 3 weeks of immunosuppression expected (>30 mg prednisone or equivalent/day)
4	 Start prednisone 1-2 mg/kg/day (or equivalent dose of methylprednisolone) If no improvement in 2–3 days, add additional/alternative immune suppressant, e.g., infliximab Provide supportive care as needed 	 Discontinue immunotherapy Continue intravenous corticosteroids Start proton pump inhibitor for GI prophylaxis Add PCP prophylaxis if more than 3 weeks of immunosuppression expected (>30 mg prednisone or equivalent/day)
		or equivalent/day) Puzanov Journal for ImmunoTherany of Cance

Puzanov Journal for ImmunoTherapy of Cancer 2017

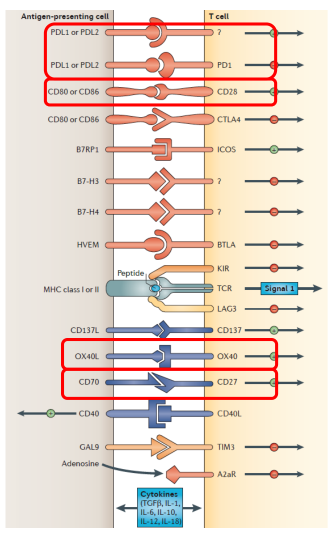








Developmental Immunotherapies for HNSCC



- Durvalumab, atezolizumab, avelumab, CK-301 (anti-PD-L1)
- Cemiplimab (anti-PD-1)
- Ipilimumab, tremelimumab (anti-CTLA-4)
- Varlilumab (anti-CD27)
- PF-04518600, tavolimab (anti-OX40)





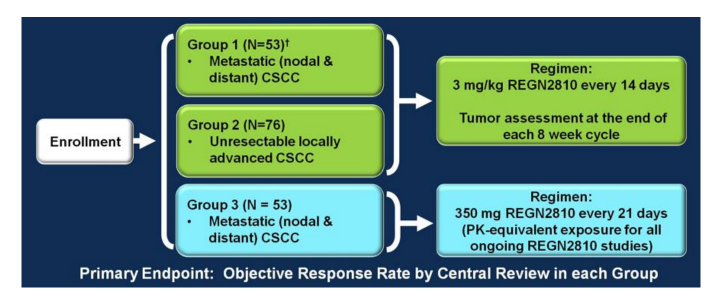




Developmental Immunotherapies for HNSCC

Cemiplimab (REGN2810) for treatment of patients with cutaneous squamous cell carcinoma (cSCC)

NCT02760498



- Largest prospective study in this disease
- ORR 46% in 82 patients in study
 - Much higher than RR in mucosal HNSCC as per KEYNOTE and CheckMate studies
- Responses durable, median DOR not reached
- Study ongoing







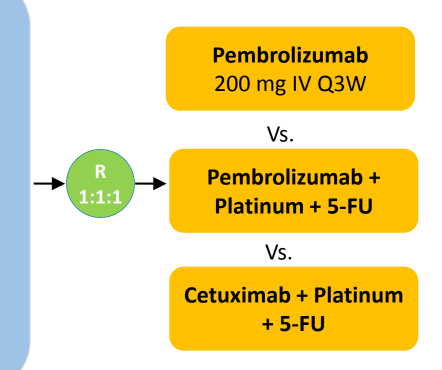


Developmental Immunotherapies for HNSCC

KEYNOTE – 048 (NCT02358031)

Key Eligibility Criteria

- R/M SCCHN of the oropharynx, oral cavity, hypopharynx, or larynx considered incurable by local therapies
- No prior systemic therapy in the R/M setting
- ECOG 0-1
- Results from HPV testing (oropharyngeal)
- Tissue for PD-L1 biomarker analysis



Primary endpoint

- PFS
- OS

Other endpoints

- PFS at 6 months
- ORR
- Biomarkers
- Quality of life









Developmental Immunotherapies for HNSCC

MASTERKEY 232/KEYNOTE-137

- Talimogene laherparepvec (T-Vec)
 - Genetically engineered herpes virus
- T-Vec 10⁶ PFU/mL <u>intratumoral injection</u> followed by 10⁸ PFU/mL Q3W
- Pembrolizumab 200 mg IV Q3W
- Eligibility:
 - R/M HNSCC not suitable for curative therapy
 - Progressed after platinum treatment
 - At least 1 injectable cutaneous, subcutaneous, or nodal tumor ≥ 10 mm in longest diameter









Patient Case Study 1

- Patient Background Information:
 - 78 yo M with a history of CAD, HTN, HLD
 - Presents with painful L sided neck mass
 - Lost 30 lbs due to anorexia



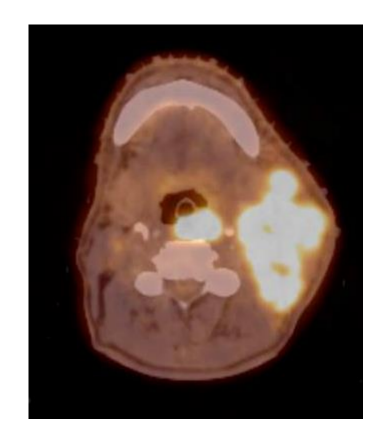






Patient Case Study 1 November 2014

- PET CT
 - Large L sided cervical mass
 - Periepiglottic tumor with no airway compromise
 - Multiple cervical osseous metastases
- Palliative hypofractionated XRT initiated







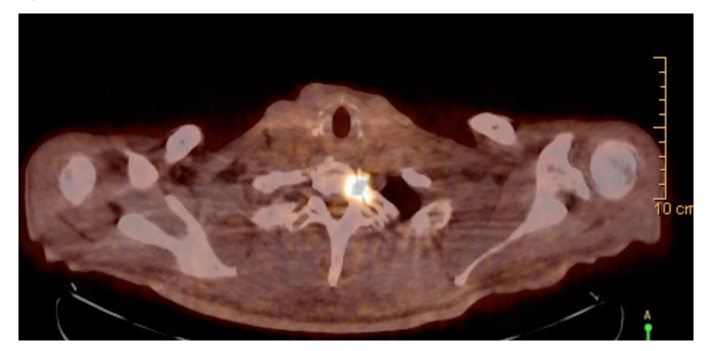




Patient Case Study 1 January 2015

- Cervical disease decreased pain improved
 - Carboplatin/paclitaxel 1st line

- PET CT revealed new osseous and axillary mets
 - Started on cetuximab 2nd line











Patient Case Study 1 June 2015

- Progression in cervical nodes
 - Re-irradiation not an option
- Enrolled in KEYNOTE-055
 - Started on pembrolizumab











Patient Case Study 1 October 2015

- Patient experienced near CR
 - Response lasted 1 year
 - No side effects of note









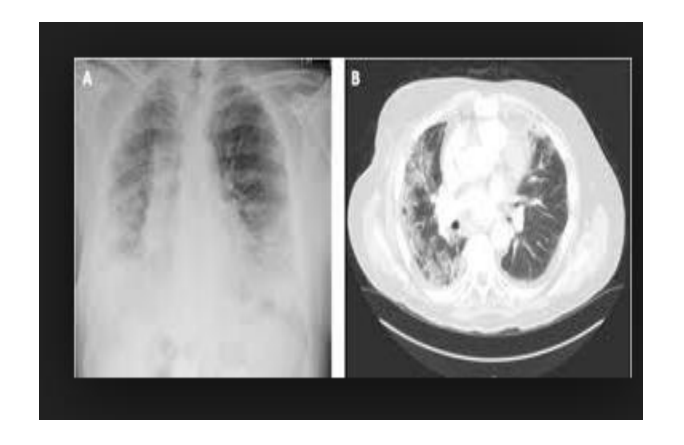


Patient Case Study 1 December 2015

Patient develops worsening
 SOB, cough, and fever to 101.5

O2 Saturation 84% on RA

PE demonstrates rales











Patient Case Study 1 December 2015

Admitted to the hospital

Started on Prednisone 1mg/kg

Symptoms improved over next
 10 days and pt back to baseline

Steroids weaned over 4 weeks











Conclusions

- 1. Chemotherapy offers short survival with many side effects
- 2. PD-1 antibodies nivolumab and pembrolizumab are approved in *platinum-refractory* recurrent / metastatic HNSCC.
- 3. Most patients have fewer side effects on PD-1 Abs than on chemotherapy
- 4. Clinical trials are underway to improve immunotherapy response rates and testing immunotherapy in other settings





