



Immunotherapy for the Treatment of Head and Neck Cancer

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Society for Immunotherapy of Cancer





- Consulting Fees: Merck Sharp and Dohme, Sanofi Genzyme, Regeneron
- Contracted Research: Boehringer Ingelheim
- I will be discussing non-FDA approved indications during my presentation.





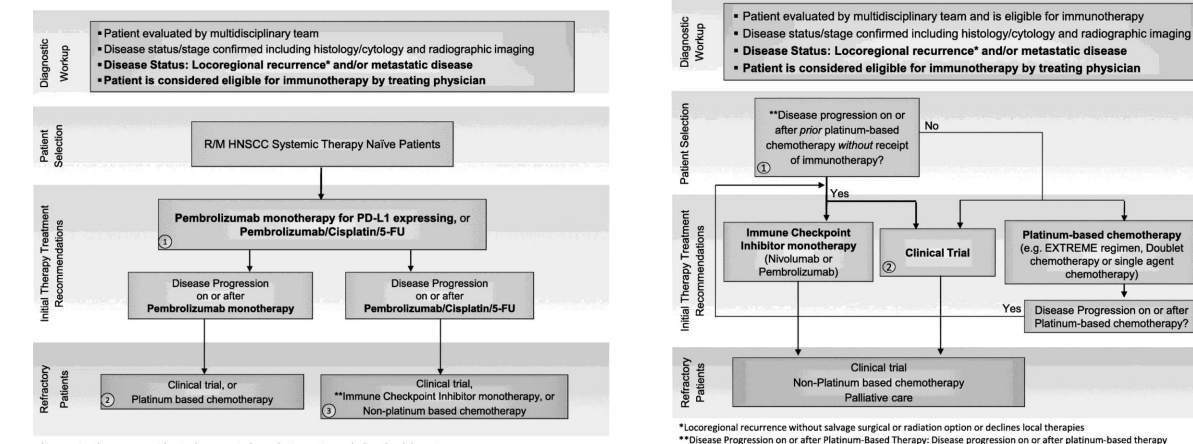


- Approved immunotherapies in head and neck cancers
- Biomarkers and immunotherapy responsiveness
- Unique considerations for head and neck cancers
- Future directions





Immunotherapy in head and neck cancer treatment



*Locoregional recurrence without salvage surgical or radiation option or declines local therapies

**Refer to Figure 2. Initial Therapy Treatment Recommendations: Immune Checkpoint Inhibitor monotherapy (nivolumab or pembrolizumab)



including within 6 months of platinum-based CRT given in the locally advanced setting. Patients that receive but cannot

Yes

Platinum-based chemotherapy

(e.g. EXTREME regimen, Doublet

chemotherapy or single agent

chemotherapy)

Disease Progression on or after Platinum-based chemotherapy?



Approved checkpoint inhibitors in head and neck cancers

Drug	Approved	Indication	Dose	
Pembrolizumab	2016	Recurrent/metastatic HNSCC, progression on/after chemotherapy	200 mg Q3W or 400 mg Q6W	
Nivolumab	2016	Recurrent/metastatic HNSCC, progression on/after chemotherapy	240 mg Q2W or 480 mg Q4W	
Pembrolizumab + platinum + fluorouracil	2019	Recurrent/metastatic HNSCC 1 st line – all patients	200 mg Q3W or 400 mg Q6W	
Pembrolizumab	2019	Recurrent/metastatic HNSCC 1^{st} line – PD-L1 CPS ≥ 1	200 mg Q3W or 400 mg Q6W	



Clinical trials in HNSCC

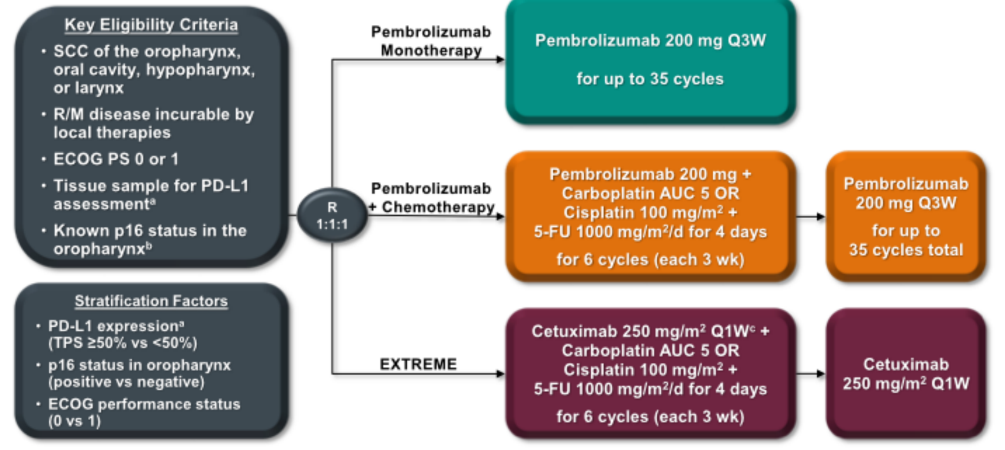
Trial	Patient selection criteria	Treatment arm(s)	Ν	ORR	Median PFS (months)	Median OS (months)
	Untreated R/M HNSCC (total population)	Pembrolizumab	301	16.9%	2.3	11.5
		Pembrolizumab + chemo	281			13.0
		Cetuximab + chemo	300	36.0%	5.2	10.7
KEYNOTE-012	R/M HNSCC	Pembrolizumab	192	18% (PD-L1+: 21%, PD-L1-: 6%)	2.1	8
CheckMate 141 R/M HNSCC with progression on platinum		Nivolumab	240	13.1% (PD-L1+: 17.7%, PD-L1-: 11.8%)	2.0	7.7
		Investigator's choice	121	5.8%	2.3	5.1
KEYNOTE-040	R/M HNSCC with progression on platinum	Pembrolizumab	247	14.6%	2.1	8.4
		Investigator's choice	248	10.1%	2.3	6.9

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KEYNOTE-048: Pembrolizumab +/-Chemotherapy in newly diagnosed R/M HNSCC



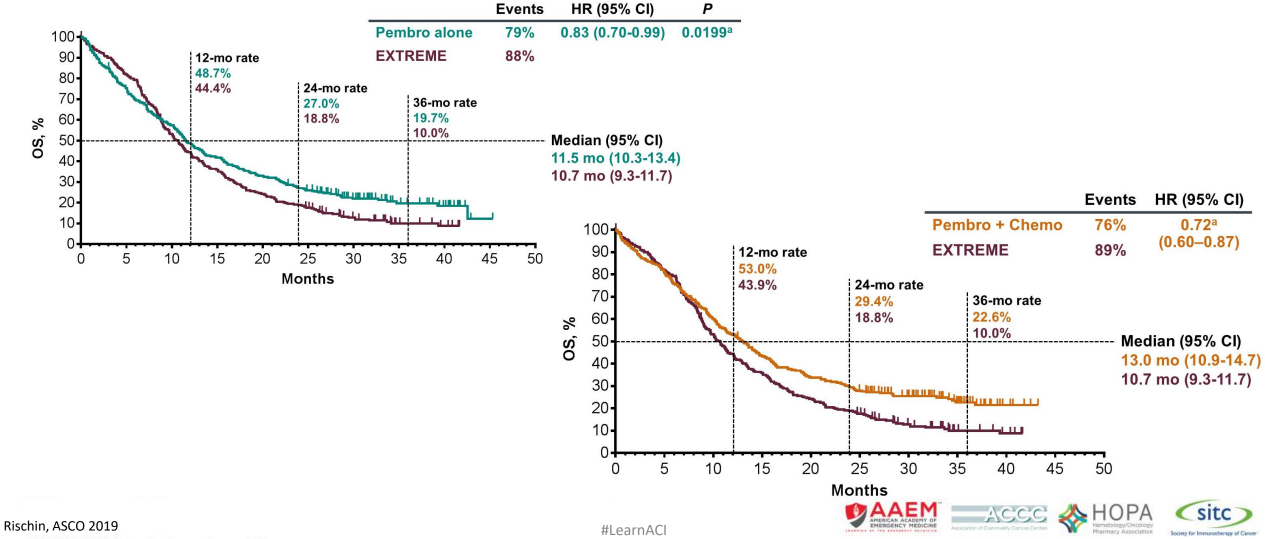
"Assessed using the PD-L1 IHC 22C3 pharmDx assay (Agilent). TPS = tumor proportion score = % of tumor cells with membranous PD-L1 expression. "Assessed using the CINtec p16 Histology assay (Ventana); cutpoint for positivity = 70%. "Following a loading dose of 400 mg/m².

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KEYNOTE-048: Overall survival in the total population

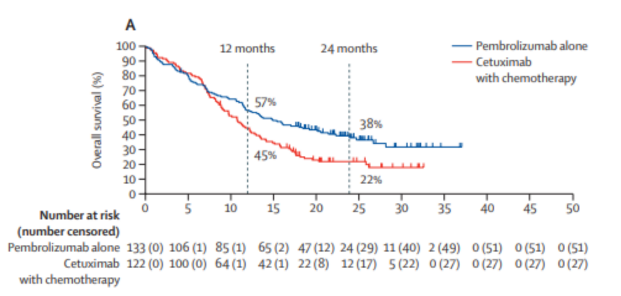


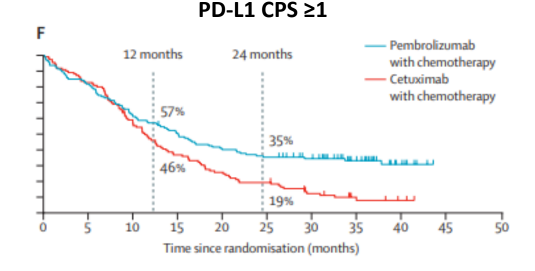
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KEYNOTE-048: Overall survival in the PD-L1 positive population

PD-L1 CPS ≥1





126 (0) 102 (0) 77 (0) 60 (1) 50 (1) 44 (1) 36 (8) 21 (22) 4 (38) 0 (42) 0 (42) 110 (0) 91 (0) 60 (1) 40 (1) 26 (1) 19 (2) 11 (4) 4 (8) 1 (11) 0 (12) 0 (12)



Burtness, Lancet 2019 © 2020–2021 Society for Immunotherapy of Cancer



KEYNOTE-048: Outcomes on subsequent therapy

Pembro **Key Eligibility Criteria** 200 mg Q3W Pembro • SCC of the oropharynx, oral cavity, hypopharynx, or larynx for up to 35 cycles R/M disease incurable by local • ECOG PS 0 or 1 Pembro Tissue sample for PD-L1 Pembro 200 mg Q3W Subsequent assessmenta + Chemo R for up to 35 PD 1:1:1 (Investigator's Known p16 status in the cycles total oropharynx^b + **Chemo**^d **Stratification Factors** • PD-L1 expression^a (TPS ≥50% vs Cetuximab

- p16 status in oropharynx (positive vs negative)
- ECOG performance status (0 vs 1)

AAEM ACCC

SITC

Therapy

choice)

<50%)

therapies

EXTREME

250 mg/m²

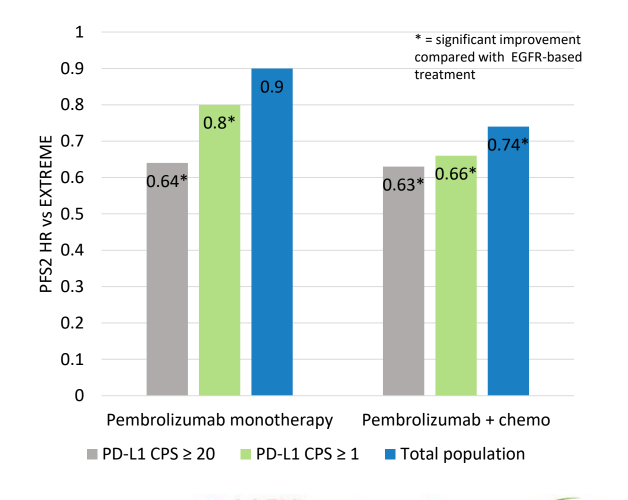
Q1W^c +

Chemod



KEYNOTE-048: Outcomes on subsequent therapy

- After progression, most common next treatment was a chemotherapy regimen
- PFS2: Progression-free survival on second treatment (after progression on KEYNOTE-048 treatment)
- Benefits seen for patients who received pembrolizumab regimens up-front
- Provides support to use of immunotherapy in front-line setting







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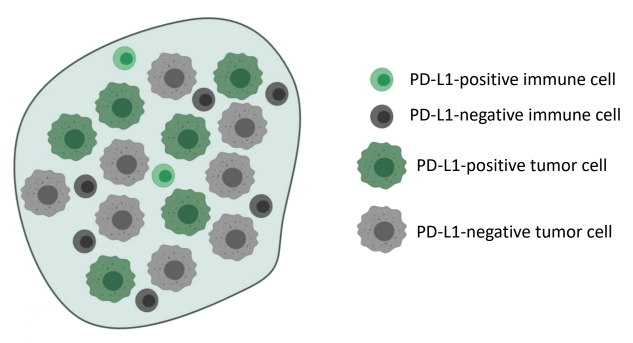




PD-L1: TPS vs CPS

 $TPS = \frac{\# of \text{ PD-L1 positive tumor cells}}{number of viable tumor cells} \times 100$

 $CPS = \frac{\# of \text{ PD-L1 positive cells (tumor cells, lymphocytes, macrophages)}}{total number of tumor and immune cells} \times 100$



$$TPS = \frac{6 \text{ positive tumor cells}}{14 \text{ total tumor cells}} \times 100 = 43$$

$$CPS = \frac{6 \text{ positive tumor cells+2 positive immune cells}}{22 \text{ total cells}} \times 100 = 36$$





Impact of PD-L1 in HNSCC

PD-L1 CPS

- KEYNOTE-048
 - First-line treatment
 - Approval of pembrolizumab monotherapy: CPS <u>></u> 1
- KEYNOTE-040
 - After platinum
 - Improved outcomes in PD-L1positive patients (by CPS > 1), no significance in total population

PD-L1 TPS

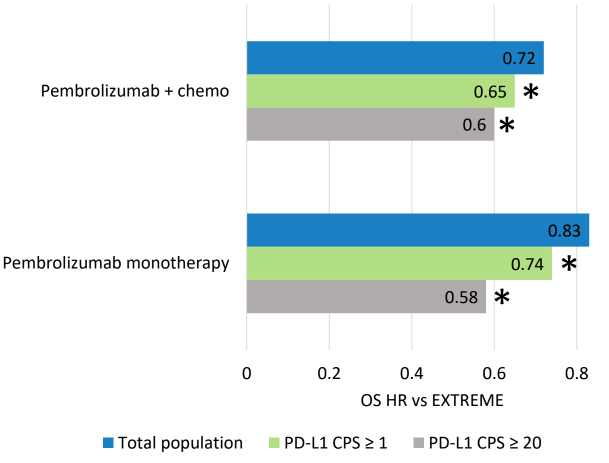
- CheckMate 141
 - After platinum
 - Greatest benefit seen for PD-L1positive tumors (TPS <u>></u> 1%), but benefit regardless
- KEYNOTE-012
 - Second-line treatment
 - Higher response rate with PD-L1 CPS-positive tumors
 - No difference for PD-L1-positive tumors by TPS





KEYNOTE-048: Outcomes by PD-L1 status

- Greatest benefits seen in tumors with highest PD-L1 expression
- Approval requires PD-L1 expression (CPS) only for monotherapy
- For total population, only pembrolizumab + chemotherapy should be considered, not monotherapy

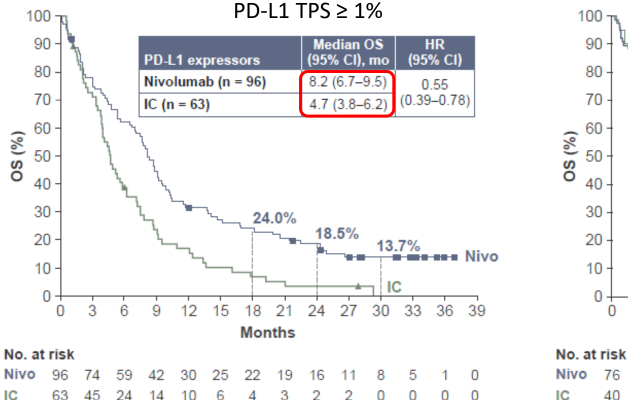


*superiority statistically demonstrated at interim or final analysis

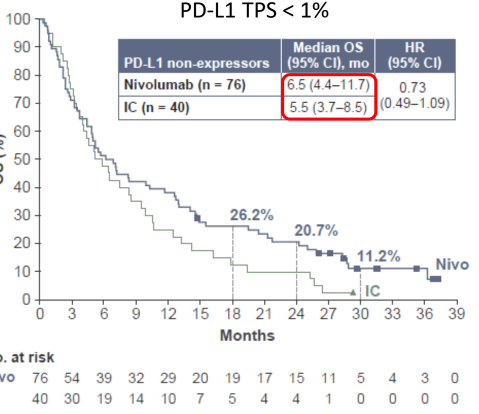




CheckMate 141: Outcomes by PD-L1 status



CheckMate 141: 2 year update



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sitc

Ferris, Oral Oncol 2018.

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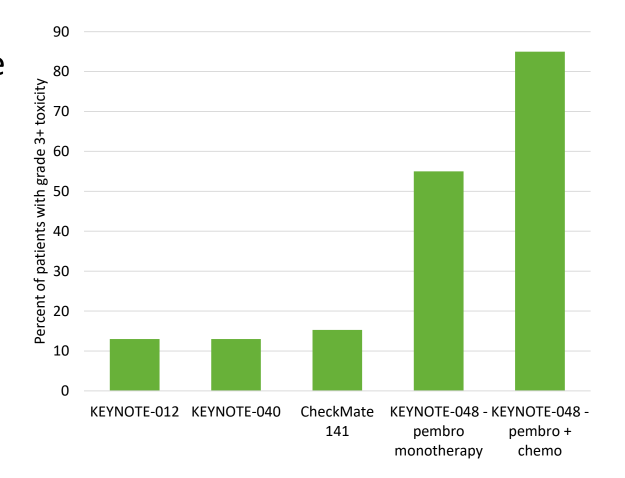
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Toxicities in head and neck cancer patients

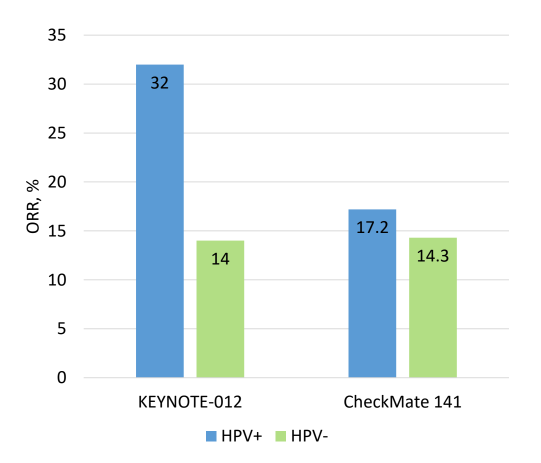
- Patients typically receive aggressive radiation treatment, with accompanying side effects
- Radiation in combination with chemotherapy, immunotherapy and/or surgery can further complicate toxicity profiles
- While combinations may have higher response rates, also have higher toxicity rates





Viral infections in HNSCC

- Virally-associated cancers are biologically and clinically distinct
 - Human papillomavirus associated with oropharynx cancer
 - Epstein Barr virus associated with nasopharyngeal cancer
- Evidence that HPV+ tumors may perform better, but there is benefit with immunotherapy regardless of HPV status







Combination immune checkpoint inhibition in HNSCC – *limited success to date*

Trial	Patient population	Treatment arms	ORR	Median OS (months)	Landmark OS
EAGLE R/M HNSCC after platinum	Durvalumab	17.9%	7.6	24-months: 18.4%	
	platinum	Durvalumab + tremelimumab	18.2%	6.5	24-months: 13.3%
		SoC	17.3%	8.3	24-months: 10.3%

Trial	Patient population	Treatment arms	Expected study completion	
KESTREL	Untreated HNSCC	Durvalumab	February 2021	
		Durvalumab + tremelimumab		
		SoC		
CheckMate 714	Platinum-refractory HNSCC	Nivolumab + ipilimumab	January 2024	
		Nivolumab		
CheckMate 651	Untreated HNSCC	Nivolumab + ipiliumumab	February 2026	
		EXTREME regimen		

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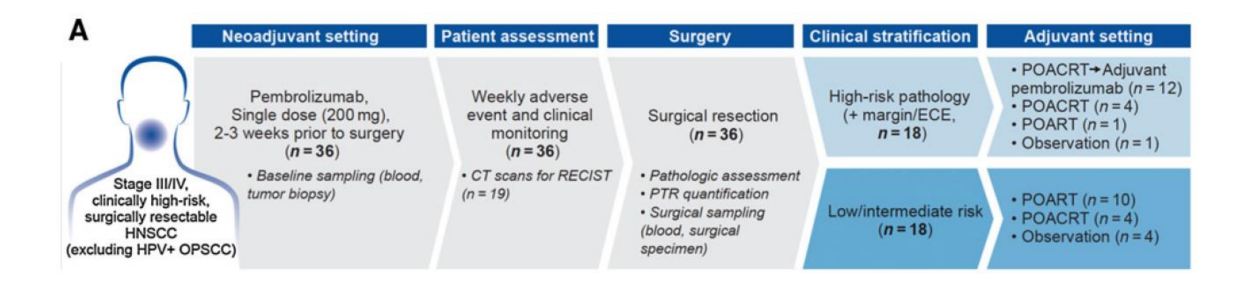


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In development: Oral cavity cancer

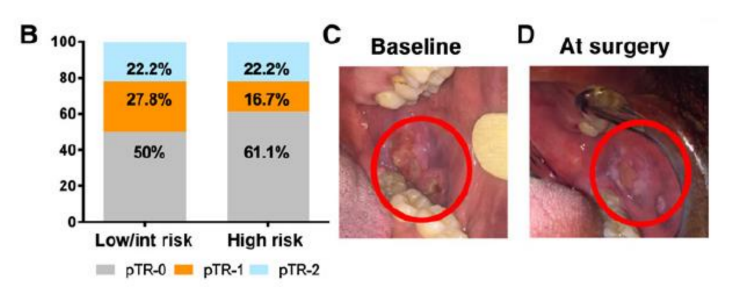






In development: Oral cavity cancer

- No serious AEs or unexpected surgical complications/delays
- pTR-2: 22%
- pTR-1: 22%
- 1-year relapse rate: 16.7%







In development: Checkpoint inhibitors + radiotherapy as primary therapy

- NCT03247712: neoadjuvant nivolumab + SBRT
 - Phase I
 - Decreased tumor size prior to surgery; high pathologic CR rate
- KEYNOTE-412: pembrolizumab + chemoradiation
 - Phase III
 - Safety confirmed, estimated completion 2021
- JAVELIN Head and Neck 100: avelumab + chemoradiation
 - Phase III trial terminated in early 2020, due to likelihood of limited efficacy
- REACH: avelumab + cetuximab + radiotherapy
 - Phase III
 - Safety confirmed, estimated completion 2027

Leidner, AACR 2019



In development: cetuximab + pembrolizumab for recurrent metastatic disease

- Cetuximab and pembrolizumab are both approved as monotherapies for HNSCC
- Phase II trial testing cetuximab + pembrolizumab:
 - Platinum refractory or ineligible disease
 - ORR: 45%
 - Median OS: 18.4 months
 - Safety profile consistent with individual drugs





In development: Selected ongoing combination trials

Trial	Patient population	Treatment arms	Targets	Expected study completion	
LEAP-010 Untreated recurrent/ metastatic PD-L1+ HNSCC (CPS ≥ 1)	metastatic PD-L1+	Pembrolizumab + lenvatinib	PD-1 + multikinase inhibitor	April 2024	
	Pembrolizumab	PD-1			
m	Untreated recurrent/ metastatic PD-L1+ HNSCC (CPS <u>></u> 1)	Pembrolizumab + GSK609	PD-1 + ICOS	July 2023	
		Pembrolizumab	PD-1		
NCT02643550	HNSCC after 1-2 therapies, including progression on Pt	Monalizumab + cetuximab	NKG2A + EGFR	Phase 1/2: 2021 Phase 3: planned	





Conclusions

- Cytotoxic chemotherapy achieves limited survival in HNSCC with unfavorable side effects.
- Checkpoint inhibitors that target the PD-1 axis, nivolumab and pembrolizumab, are approved in platinum-refractory/exposed recurrent/metastatic HNSCC.
- Nivolumab and pembrolizumab are in general better tolerated than cytotoxic chemotherapy.
- Ongoing areas of research include combinations of immunotherapy with radiation and/or other drugs and development of predictive biomarkers.









Cohen et al. Journal for ImmunoTherapy of Cancer (2019) 7:184 https://doi.org/10.1186/s40425-019-0662-5

Journal for ImmunoTherapy of Cancer

POSITION ARTICLE AND GUIDELINES

The Society for Immunotherapy of Cancer consensus statement on immunotherapy for the treatment of squamous cell carcinoma of the head and neck (HNSCC)

Ezra E. W. Cohen¹, R. Bryan Bell², Carlo B. Bifulco², Barbara Burtness³, Maura L. Gillison⁴, Kevin J. Harrington⁵, Quynh-Thu Le⁶, Nancy Y. Lee⁷, Rom Leidner², Rebecca L. Lewis⁸, Lisa Licitra⁹, Hisham Mehanna¹⁰, Loren K. Mell¹, Adam Raben¹¹, Andrew G. Sikora¹², Ravindra Uppaluri¹³, Fernanda Whitworth¹⁴, Dan P. Zandberg⁸ and Robert L. Ferris^{8*}



Open Access







Case Studies









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- 1. A 48yoM with HPV+ locally-advanced left tonsil cancer treated with definitive chemoradiation 2 years ago, presents to the ED with a new left neck lump. CT Neck and Chest is obtained in the ED and shows 2 enlarged nodes in the left neck and multiple new lung nodules, 1-2 cm in size, concerning for recurrent and metastatic disease. You are consulted as the on-call oncologist. What would you recommend next?
 - A. Admit for inpatient evaluation
 - B. Outpatient biopsy of a lung nodule to confirm diagnosis
 - C. ENT consult for left neck dissection





- A. Admit for inpatient evaluation
 Not necessary because the patient does not seem to have an urgent indication to admit. Work-up can be completed as outpatient.
- **B.** Outpatient biopsy of a lung nodule to confirm diagnosis Most appropriate option
- C. ENT consult for left neck dissection

If he had locoregionally recurrent disease, would have been appropriate to refer to ENT for salvage neck dissection to attempt cure. But if he has metastatic disease, he needs systemic therapy and would not benefit from salvage surgery.





CT guided biopsy of a peripheral lung nodule confirms HPV+ squamous cell cancer, consistent with distant metastasis from the head and neck primary. What additional testing would you request that would help you choose first-line systemic therapy?

- A. HIV testing on the patient
- B. PD-L1 CPS on tumor biopsy
- C. Hepatitis serologies
- D. Her-2 testing on tumor biopsy







- A. HIV testing on the patientDoes not help guide systemic therapy
- **B.** PD-L1 CPS on tumor biopsy

If he has PD-L1 CPS > 1, he would be eligible for pembrolizumab monotherapy or in combination with platinum doublet

- C. Hepatitis serologies Does not help guide management
- D. Her-2 testing on tumor biopsy
 Her-2 positivity is not routinely seen in head and neck cancers







Mr. VB is a 54yoM with metastatic SCCHN. He is currently undergoing treatment with pembrolizumab monotherapy and has received 9 cycles of treatment. He presents to the ED with severe fatigue and has orthostatic hypotension. On blood work, Na level is 126. ED gives him a liter of NS and calls you as the on-call oncologist for clearance for discharge home. What would you recommend?

- A. Administer another 500cc of NS
- B. Can discharge home with outpatient f/u with primary oncologist within a week
- C. Admit for work-up





- A. Administer another 500cc of NS Although he does need continued hydration, he should undergo further work-up for adrenal insufficiency which is a life-threatening condition
- B. Can discharge home with outpatient f/u with primary oncologist within a week He should be admitted for expedited work-up

C. Admit for work-up

His Fatigue, hypotension and hyponatremia may be indicative of adrenal insufficiency. This should be suspected and worked up in this patient who has been on immune checkpoint inhibition for several weeks





You ask the ED to obtain SIADH labs, serum TSH, ACTH and cortisol levels. Work-up is negative for SIADH, TSH is 7, ACTH is high and cortisol is 1. What would you recommend next?

- A. Cosyntropin stim test and start hydrocortisone
- B. Obtain MRI brain to evaluate for hypophysitis
- C. Discharge home after another liter of NS if orthostatic hypotension is resolved and Na levels are improved





A. Cosyntropin stim test and start hydrocortisone

His clinical presentation and low cortisol/high ACTH are very suggestive of primary adrenal insufficiency secondary to pembrolizumab induced adrenalitis. This can be confirmed by an inadequate response to the cosyntropin stimulation test and treatment should be started right away with steroids

- B. Obtain MRI brain to evaluate for hypophysitis Although this can be considered to evaluate for secondary adrenal insufficiency, the elevated ACTH and TSH support an intact pituitary axis
- C. Discharge home after another liter of NS if orthostatic hypotension is resolved and Na levels are improved Not starting steroids right away could be life-threatening for this patient. He should therefore be admitted for monitoring until work-up is completed and Na levels improve

