

## Immunotherapy for the Treatment of Head and Neck Cancer

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### **Disclosures**

- No relevant financial relationships to disclose
- I will be discussing non-FDA approved indications during my presentation.











#### Outline

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



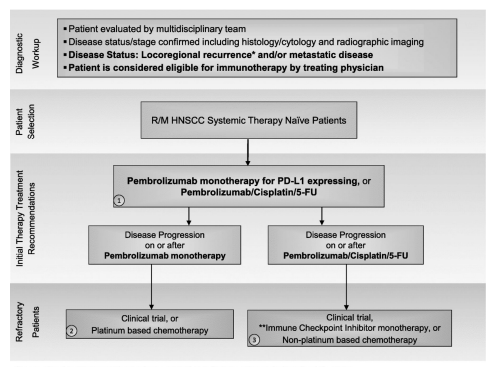








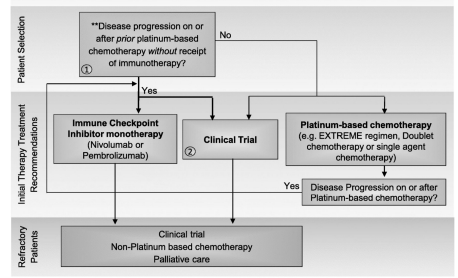
### Immunotherapy in head and neck cancer treatment



<sup>\*</sup>Locoregional recurrence without salvage surgical or radiation option or declines local therapies

ojagnostic Workup

- Patient evaluated by multidisciplinary team and is eligible for immunotherapy
- Disease status/stage confirmed including histology/cytology and radiographic imaging
- Disease Status: Locoregional recurrence\* and/or metastatic disease
- Patient is considered eligible for immunotherapy by treating physician



<sup>\*</sup>Locoregional recurrence without salvage surgical or radiation option or declines local therapies

<sup>\*\*</sup>Disease Progression on or after Platinum-Based Therapy: Disease progression on or after platinum-based therapy including within 6 months of platinum-based CRT given in the locally advanced setting. Patients that receive but cannot tolerate platinum-based chemotherapy would also be included in this category.

HNSCC: head and neck squamous cell carcinoma









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



### 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 <sup>st</sup> line – all patients	200 mg Q3W or 400 mg Q6W	
Pembrolizumab	2019	Recurrent/metastatic HNSCC 1 <sup>st</sup> line − PD-L1 CPS ≥ 1	200 mg Q3W or 400 mg Q6W	













### Clinical trials in HNSCC

Trial	Patient selection criteria	Treatment arm(s)	N	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
	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







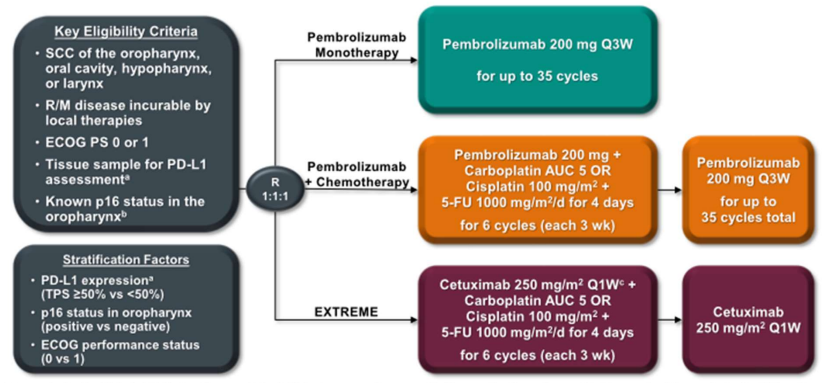


I cannot access the full KEYNOTE-048 paper - can you add in the response rate for pembro+chemo in the total population?

Emily Ehlerding, 8/31/2020



# 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.



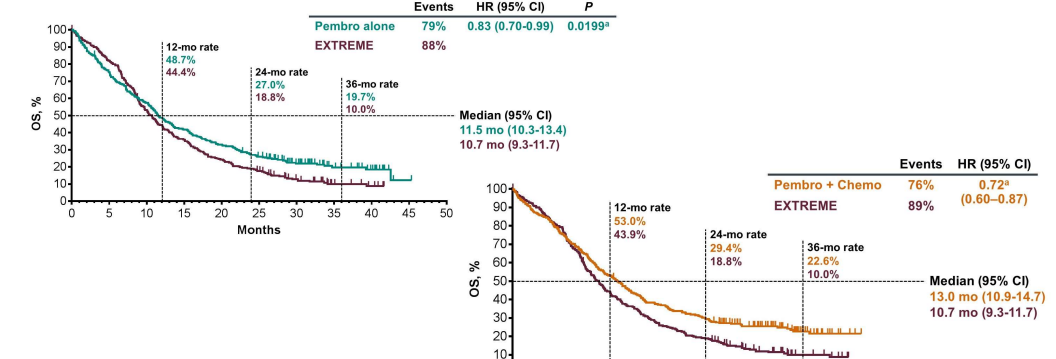








### KEYNOTE-048: Overall survival in the total population



Rischin, ASCO 2019

#LearnACI

5

10

15



40

25

**Months** 

30

20

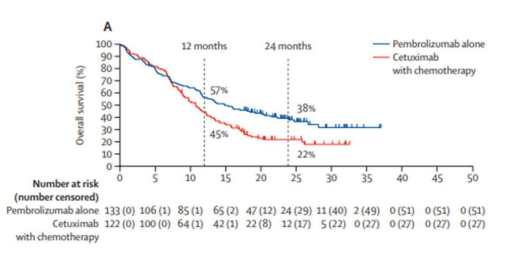


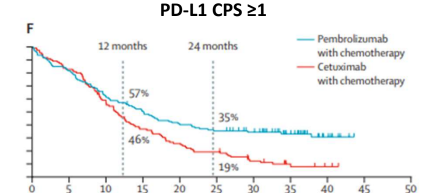




## KEYNOTE-048: Overall survival in the PD-L1 positive population

#### 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)

Time since randomisation (months)











## 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 therapies ECOG PS 0 or 1 Pembro · Tissue sample for PD-L1 Pembro 200 mg Q3W Subsequent assessment<sup>a</sup> R + Chemo for up to 35 Therapy · Known p16 status in the 1:1:1 (Investigator's cycles total oropharynx<sup>b</sup> choice) Chemo<sup>d</sup> **Stratification Factors** • PD-L1 expressiona (TPS ≥50% vs Cetuximab <50%) **EXTREME** 250 mg/m<sup>2</sup> • p16 status in oropharynx (positive Q1Wc + vs negative) Chemo<sup>d</sup> · ECOG performance status (0 vs 1)





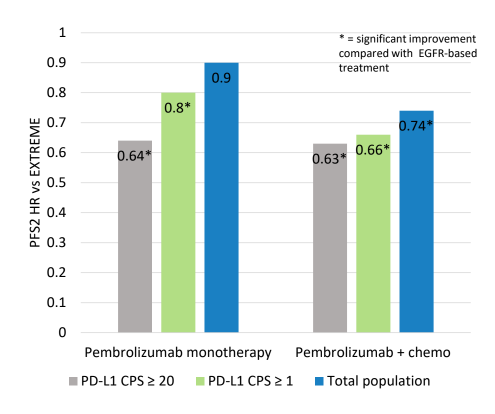






## KEYNOTE-048: Outcomes on subsequent therapy

- 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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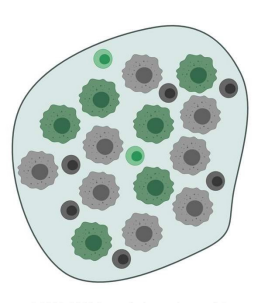




#### PD-L1: TPS vs CPS

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

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



- PD-L1-positive immune cell
- PD-L1-negative immune cell
- PD-L1-positive tumor cell
- PD-L1-negative tumor cell

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

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











### Impact of PD-L1 in HNSCC

#### PD-L1 CPS

- KEYNOTE-048
  - First-line treatment
  - Approval of pembrolizumab monotherapy: CPS > 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 ≥ 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





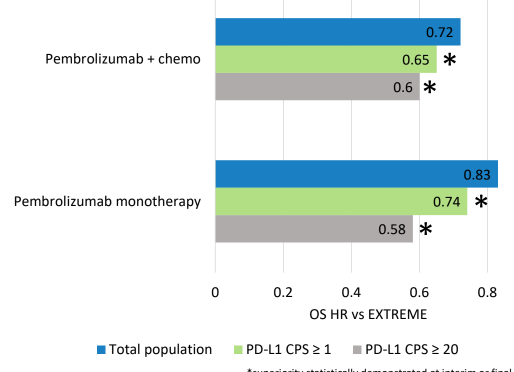






### KEYNOTE-048: Outcomes by PD-L1 status

- 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





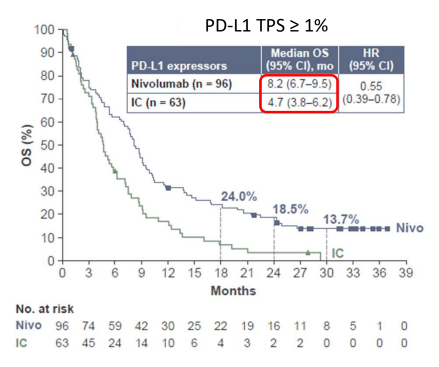


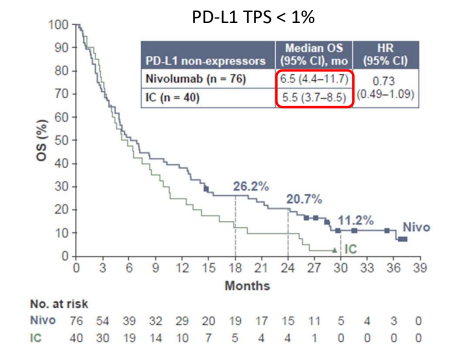




### CheckMate 141: Outcomes by PD-L1 status

#### CheckMate 141: 2 year update















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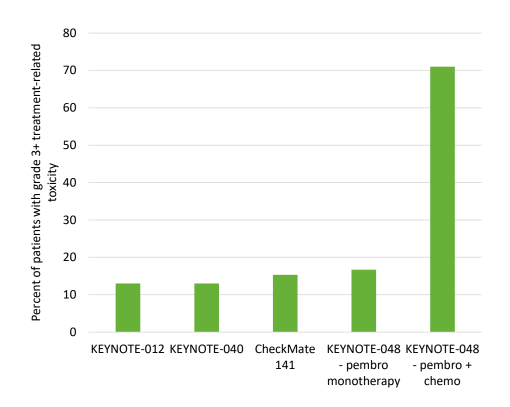






## Toxicities in head and neck cancer patients

- 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







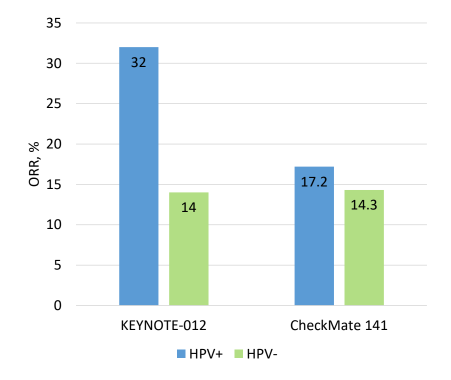






#### Viral infections in HNSCC

- 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%	
	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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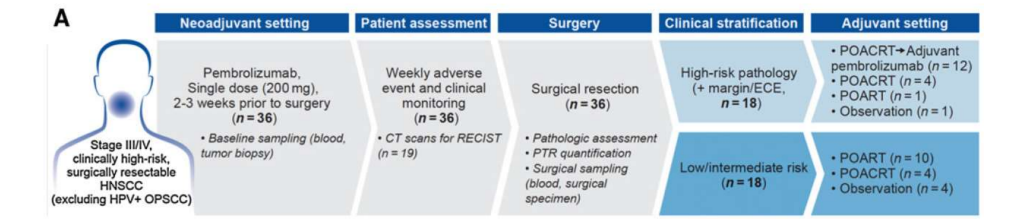








### In development: Oral cavity cancer













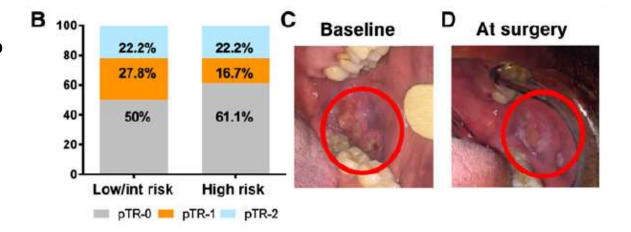
### In development: Oral cavity cancer

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











### 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)	Pembrolizumab + lenvatinib	PD-1 + multikinase inhibitor	April 2024	
		Pembrolizumab	PD-1		
INDUCE-3 Untreated recurrent metastatic PD-L1+ HNSCC (CPS ≥ 1)		Pembrolizumab + GSK609	PD-1 + ICOS	July 2023	
	HNSCC (CPS $\geq$ 1)	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.











#### Resources



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

Journal for ImmunoTherapy of Cancer

#### **POSITION ARTICLE AND GUIDELINES**

**Open Access** 

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. Cohen<sup>1</sup>, R. Bryan Bell<sup>2</sup>, Carlo B. Bifulco<sup>2</sup>, Barbara Burtness<sup>3</sup>, Maura L. Gillison<sup>4</sup>, Kevin J. Harrington<sup>5</sup>, Quynh-Thu Le<sup>6</sup>, Nancy Y. Lee<sup>7</sup>, Rom Leidner<sup>2</sup>, Rebecca L. Lewis<sup>8</sup>, Lisa Licitra<sup>9</sup>, Hisham Mehanna<sup>10</sup>, Loren K. Mell<sup>1</sup>, Adam Raben<sup>11</sup>, Andrew G. Sikora<sup>12</sup>, Ravindra Uppaluri<sup>13</sup>, Fernanda Whitworth<sup>14</sup>, Dan P. Zandberg<sup>8</sup> and Robert L. Ferris<sup>8\*</sup>







