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PHASE II EVALUATION OF THE TRIPLE COMBINATION OF PDS0101, M9241, AND BINTRAFUSP ALFA IN PATIENTS WITH HPV 16 POSITIVE MALIGNANCIES

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Introduction

- >630,000 new cases of HPV-related cancer (e.g. cervical, oropharyngeal, anal) reported worldwide annually¹
- PD-1 inhibitors have been evaluated in these tumor types and ORRs have ranged from 13–24%²⁻⁸
- Nivolumab & pembrolizumab are FDA approved for HNSCC; pembrolizumab is approved for PD-L1+ cervical cancer
- Unfortunately, the majority of patients who receive these anti PD-1 inhibitors will progress
- For these patients with checkpoint refractory disease there is no clear effective standard of care therapy
- HPV infection is also linked to upregulation of TGF-β signaling⁹ - Genome-wide association studies showed that TGF-βR1 is significantly overexpressed in HPV-related cancer¹⁰

1. de Martel C, et al. Int J Cancer. 2017;141:664–70; 2. Viens LJ, et al. MMWR Morb Mortal Wkly Rep.; 2. Bauml J, et al. J Clin Oncol 2017;35:1542–49; 3. Ott PA, et al. Ann Oncol. 2017;28:1036–41; 4. Hollebecque A, et al. J Clin Oncol. 2017;35(Suppl):Abstract 5504; 5. Chung HC, et al. J Clin Oncol. 2018;36(Suppl):Abstract 5522; 6. Ferris RL, et al. N Engl J Med. 2016;375:1856–67; 7 Mehra R, et al. Br J Cancer. 2018;119:153–59; 8 Morris VK, et al. Lancet Oncol. 2017;18:446–53. 2016;65:661–66; 9. Torres-Poveda K, et al. World J Clin Oncol. 2014;5:753-63; 10. Levovitz C, et al. Cancer Res. 2014;74:6833-44.













Bintrafusp alfa: a TGF-ß and PD-L1 Inhibitor



1.Strauss J, et al. Clin Cancer Res. 2018;24:1287–95; 2. Paz-Ares L, et al. J Clin Oncol. 2018;36(Suppl):Abstract 9017; 3. Cho BC, et al. Ann Oncol. 2018;29(Suppl):Abstract 10480.

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- Bintrafusp alfa is an innovative • first-in-class bifunctional fusion protein composed of the extracellular domain of the **TGF-**βRII receptor (a TGF-β "trap") fused to a human IgG1 mAb blocking PD-L1
- In a phase 1 study, bintrafusp alfa was well • tolerated and produced durable responses in several solid tumor types ¹⁻³







Bintrafusp alfa in HPV-Related Cancers



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- 79 patients with advanced HPV-associated cancers ullet(59 checkpoint naïve and 20 checkpoint refractory) received bintrafusp alfa IV every 2 weeks until disease progression or intolerance¹
- Side effect profile similar to standard anti-PD(L)1 inhibitors with the addition of keratoacanthomas & mucosal bleeding
- ORR was 30.5% in checkpoint naïve disease
- ORR was 10% in checkpoint refractory disease ullet







PDS0101: Versamune[®]-based HPV 16 cancer vaccine



1.Wood LV, et al., SITC 2019, (O19) Abstract ID 12533



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- Micellar multi-peptide based therapeutic vaccine targeting HPV 16 E6/E7 (HPV 16 is the genotype responsible for majority of HPVrelated cancers worldwide)
- Versamune[®] nanoparticles contain the cationic lipid R-DOTAP which upregulates type I IFNs and promotes antigen cross-presentation
- In a phase I trial patients with cervical intraepithelial neoplasia developed strong HPV-specific CD4+ and CD8+ T cell immune responses¹
- Was well tolerated with mild transient site reactions and minimal systemic toxicity









NHS-IL12 Immunocytokine. (A) NHS76 is a fully human 2nd generation TNT antibody bound to 2 murine IL-12 (p70) molecules. (B) a: Specific tumor targeting of transplanted lung carcinoma by the MAb NHS-IL12(mu). Control MAb BC1-IL12(mu). b: NHS-IL12 tumor targeting of nuclear DNA histones.

1.Strauss J, et al. Clin Cancer Res. 2019 Jan 1;25(1):99-109

M9241 (NHSIL12)

- Tumor targeting IL12 immunocytokine
- Composed of two IL12 heterodimers fused to NHS76 antibody which binds to histones on free DNA fragments found in areas of tumor necrosis
- In phase 1 trial in patients with advanced solid tumors the most frequently observed AEs included flu like symptoms and asymptomatic lab abnormalities (e.g. mild cytopenias and liver enzyme elevations)¹
- M9241 treatment resulted in increased T cell infiltration in the TME

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Freelinical Combo of PDS0101, M9241 & Bintrafusp alfa



- \bullet
- ulletinfiltration in the TME and tumor reduction¹

1.Rumfield C, J Immunother Cancer. 2020 Jun;8(1):e000612

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Single, double and triple combinations were evaluated in a TC-1 HPV16+ tumor model Triple combination generated the maximum HPV-specific immune response, T cell





Study Design

- wks'[NCT04287868]



Treatment until confirmed progression, unacceptable toxicity, or any criteria for withdrawal; treatment past progression was allowed

Patients with advanced HPV-related cancers received the combination of bintrafusp alfa at 1200 mg flat dose i.v. q 2wks, M9241 at 16.8 mcg/kg s.c. q 4 wks and PDS0101 given as two separate 0.5 ml s.c. injections q 4

Dose reductions of M9241 to 8 mcg/kg were allowed as well as skipped doses of agent(s) for toxicities

HPV genotyping was done with PCR based assays (BD Onclarity or Molecular MD) if testing not already done











	All patients N=25
Age, median (range), years	50 (37-80)
Female, n (%)	17 (68)
Tumor type, n (%) Cervical Anal Head & Neck SCC Vulvar/ Vaginal	10 (40) 6 (24) 6 (24) 3 (12)
Number of prior anticancer therapies, n (%) 1 2 ≥3	5 (20) 11 (44) 9 (36)
Prior chemotherapy, n (%)	25 (100)
Prior radiotherapy, n (%)	24 (96)
Prior PD-(L)1 inhibitor therapy, n (%)	14 (56)
HPV status, n (%) HPV 16 HPV type other than 16 Negative	18 (72) 6 (24) 1 (4)

Key baseline patient and disease characteristics

- As of 01 MAR 2021, 25 patients had received the \bullet triple combination of PDS0101, M9241 & bintrafusp alfa
 - The median follow-up is 8 months





	N=25
	Grade ≥2
Treatment-related adverse events (TRAEs)	23 (92)
TRAEs leading to discontinuation of ≥ 1 drug(s)	5 (20)
Treatment-related serious AEs	7 (28)
TRAEs in ≥5% of patients	
Anemia	12 (48)
Lymphocyte decrease	7 (28)
Flu like symptoms	6 (24)
Injection site reactions	5 (20)
Hematuria	4 (16)
AST/ ALT/ Alk phos elevation	4 (16)
Keratoacanthomas	4 (16)
Leukocyte decrease	3 (12)
Maculopapular rash	3 (12)
Pruritis	3 (12)
Nausea/ vomiting	3 (12)
Mucositis	3 (12)
Hypothyroidism	3 (12)
Peripheral motor neuropathy	2 (8)
Fatigue	2 (8)

1. Hemophagocytic lymphohistiocytosis

Safety summary

- Grade 3 TRAEs occurred in 10 (40%) patients
 - anemia due to hematuria (n=4), AST/ALT elevation (n=2); flu like symptoms (n=1), nausea/ vomiting (n=1), leukopenia (n=1), lymphopenia (n=2), HLH¹ (n=1)
- All four patients with grade 3 hematuria had cervical ca with prior pelvic RT + brachytherapy
- One patient with transient grade 3 leukopenia and lymphopenia also had transient grade 4 neutropenia
- 4 patients who originally had grade 3 toxicities with the triple combo including M9241 at 16.8 mcg/kg tolerated the triple combo with M9241 at 8 mcg/kg w/o any further grade ≥3 toxicities
- No treatment-related deaths occurred

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	All patient s N=25	HPV 16+ N=18	HPV 16+ CPI Naïve N=6	HPV 16+ CPI Refractory N=12
BOR, n (%) Complete response (CR) Partial response (PR)	2 (8) 8 (32)	2 (11.1) 8 (44.4)	1 (16.7) 4 (66.7)	1 (8.3) 4 (33.3)
ORR (CR+PR), n (%)	10 (40)	10 (55.6)	5 (83.3)	5 (41.7)
Disease Reduction, n (%)	13 (52)	12 (66.7)	5 (83.3)	7 (58.3)
Ongoing response, n/n (%)	8/10 (80)	8/10 (80%)	4/5 (80%)	4/5 (80%)
Overall Survival, n/n (%)*	20/25 (80)	16/18 (88.9)	6/6 (100)	10/12 (83.3)

* Median 8 months of follow up

1. Bauml J, et al. J Clin Oncol 2017;35:1542–49; 2. Ott PA, et al. Ann Oncol. 2017;28:1036–41; 3. Mehra R, et al. Br J Cancer. 2018;119:153–59; 4. Ferris RL, et al. N Engl J Med. 2016;375:1856–67; 5. Morris VK, et al. Lancet Oncol. 2017;18:446–53; 6. Chung HC, et al. J Clin Oncol 2019;37: 1470-8; 7. Strauss J, et al. J Immunother Cancer. 2020 Dec;8(2):e001395

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Patient Outcomes

- ORR 55.6% (tumor reduction 66.7%) in HPV 16+ disease
- ORR 83.3% in CPI <u>naïve</u> HPV 16+ disease
- ORR 41.7% (tumor reduction 58.3%) in CPI <u>refractory</u> HPV 16+ disease
- After a median 8 months of follow up:
 - 80% of responses are ongoing ullet
 - 6/6 (100%) pts with HPV 16+ CPI naïve disease ulletremain alive (historical median OS is 7-11 mo)¹⁻⁶
 - 10/12 (83.3%) pts with HPV 16+ CPI refractory ulletdisease remain alive (historical median OS is 3-4 mo)⁷







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100% 80% 60% 40% 20% 0% -20% -40% -60% -80% -100%

Best Overall Response

o Cervical o Vaginal/Vulvar

o Anal **o** HNSCC

• Responses in HPV 16+ disease occurred irrespective of tumor type















Overwhelming majority of HPV 16+ CPI naive pts had a response

Majority of HPV 16+ CPI refractory pts had tumor shrinkage ightarrow

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Best Overall Response

• HPV 16 CPI Primary Refractory o HPV 16 CPI Secondary Refractory

Primary Refractory: Prior PD or SD < 6 months Secondary Refractory: Prior PR or SD > 6 months





Conclusions

- Triple combination of PDS0101, M9241 and bintrafusp alfa appears to have a advanced HPV 16+ malignancies
- Clinical activity noted irrespective of tumor type or CPI status •
- •
- ORR was 83.3% in patients with CPI <u>naive</u> HPV 16+ disease •
- After a median 8 months of follow up:
 - 80% of responses are ongoing
 - 6/6 (100%) pts with HPV 16+ CPI naïve disease remain alive •
 - 10/12 (83.3%) pts with HPV 16+ CPI refractory disease remain alive
- Accrual is ongoing to the triple combination [NCT04287868] \bullet

manageable safety profile along with early evidence of notable clinical activity for pts with

ORR was 55.6% (tumor reduction 66.7%) in all pts with advanced HPV 16+ disease ORR was 41.7% (tumor reduction 58.3%) in patients with CPI <u>refractory</u> HPV 16+ disease







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Disclosures

Will be added by ASCO



