

Rapid Clearance Profile of Plasma Circulating Tumor HPV Type 16 DNA during Chemoradiotherapy Correlates with Disease Control in HPV-Associated Oropharyngeal Cancer



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Abstract

Purpose: To identify a profile of circulating tumor human papilloma virus (HPV) DNA (ctHPVDNA) clearance kinetics that is associated with disease control after chemoradiotherapy (CRT) for HPV-associated oropharyngeal squamous cell carcinoma (OPSCC).

Experimental Design: A multi-institutional prospective biomarker trial was conducted in 103 patients with (i) p16-positive OPSCC, (ii) M0 disease, and (iii) receipt of definitive CRT. Blood specimens were collected at baseline, weekly during CRT, and at follow-up visits. Optimized multianalyte digital PCR assays were used to quantify ctHPVDNA (types 16/18/31/33/35) in plasma. A control cohort of 55 healthy volunteers and 60 patients with non-HPV-associated malignancy was also analyzed.

Results: Baseline plasma ctHPVDNA had high specificity (97%) and high sensitivity (89%) for detecting newly diag-

nosed HPV-associated OPSCC. Pretreatment ctHPV16DNA copy number correlated with disease burden, tumor HPV copy number, and HPV integration status. We define a ctHPV16DNA favorable clearance profile as having high baseline copy number (>200 copies/mL) and >95% clearance of ctHPV16DNA by day 28 of CRT. Nineteen of 67 evaluable patients had a ctHPV16DNA favorable clearance profile, and none had persistent or recurrent regional disease after CRT. In contrast, patients with adverse clinical risk factors (T4 or >10 pack years) and an unfavorable ctHPV16DNA clearance profile had a 35% actuarial rate of persistent or recurrent regional disease after CRT ($P = 0.0049$).

Conclusions: A rapid clearance profile of ctHPVDNA may predict likelihood of disease control in patients with HPV-associated OPSCC patients treated with definitive CRT and may be useful in selecting patients for deintensified therapy.

Introduction

The discovery of human papilloma virus (HPV) infection as a driver of oropharyngeal carcinogenesis has been a major advancement in head and neck oncology. Patients with HPV-associated oropharyngeal squamous cell carcinoma (OPSCC) have a significantly better prognosis than patients with HPV-negative OPSCC (1). The 3-year local regional control and overall survival for HPV-associated versus HPV-negative are respectively 86% versus 65% and 82% versus 57% (1).

Further risk stratification of patients with HPV-associated OPSCC is done using tobacco smoking history: low risk (HPV-associated, ≤ 10 pack years, overall survival 93%) and intermediate risk (HPV-associated, >10 pack years, overall survival 71%; ref. 1).

It is now standard clinical practice to use HPV, smoking status, and disease stage to risk stratify patients to facilitate prognostication, and investigation of deintensified treatment regimens is an active area of clinical trial research (2, 3). Though the primary

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Translational Relevance

Plasma circulating tumor human papilloma virus (HPV) DNA (ctHPVDNA) is a promising biomarker for monitoring treatment responses in patients with HPV-associated oropharyngeal squamous cell carcinoma. This study characterizes baseline levels and clearance kinetics of ctHPVDNA in a cohort of patients treated with definitive chemoradiotherapy (CRT). A favorable clearance profile was defined as high baseline ctHPVDNA levels (>200 copies/mL) that are rapidly eliminated during CRT (>95% clearance by day 28). Patients with a favorable clearance profile, regardless of tobacco pack years, had excellent cancer control [regional disease-free survival (DFS) = 100%], whereas those with unfavorable clearance profile and adverse clinical factors (T4, >10 tobacco pack years) had significantly worse cancer control (regional DFS = 65%). Future studies assessing ctHPVDNA as an integral biomarker to guide treatment deintensification are warranted and may facilitate personalized treatment decisions based on tumor biology in addition to clinical risk factors.

biomarker of HPV status and secondary modifier of tobacco pack years (TPY) allow for significant prognostication, they are imprecise. For example, >10 TPY did not affect cancer control in HPV-associated OPSCC patients in a large retrospective study (4). Others have also reported lower tumor HPV copy number, presence of non-HPV16 viral strains, and HPV integration as being associated with adverse prognosis (5–12). Thus, there is a need to identify additional biomarkers of therapeutic sensitivity that may help to better risk stratify patients.

Circulating tumor DNA (ctDNA) is released by dying cancer cells and represents an accessible source for detecting tumor genetic biomarkers in many cancer types, including OPSCC (13). Plasma ctDNA can be evaluated serially over time to analyze the profile of molecular ctDNA responses during cancer therapy (14). The clinical utility of Epstein–Barr Virus (EBV) ctDNA has been established for the early detection (15) of EBV-associated nasopharyngeal cancer and prognostication of patients after definitive chemoradiotherapy (CRT; ref. 16). Similarly, plasma circulating tumor HPV16 DNA (ctHPV16DNA) is detectable in the majority of patients with HPV-associated OPSCC (12, 17–20). Limited prior data have shown that ctHPV16DNA levels become largely undetectable after CRT in most patients, and that ctHPV16DNA levels may increase at the time of disease recurrence (12, 19–21). These prior studies retrospectively analyzed limited subsets of patients using real-time PCR.

We have designed and validated a highly sensitive and specific digital PCR (dPCR) assay for absolute quantification of ctHPV16DNA, as well as circulating tumor HPV DNA of the four most prevalent alternative high-risk HPV strains (18/31/33/35). We integrated these ctHPVDNA analyses into our ongoing multi-institutional prospective phase II clinical trials for patients with HPV-associated OPSCC, to investigate whether ctHPVDNA baseline levels and clearance kinetics improve risk stratification when combined with established clinical risk factors. We herein report our observations regarding a ctHPV16DNA rapid clearance profile that, in combination with established clinical risk factors, predicts local and regional disease control after definitive CRT.

Materials and Methods

Ethics approval and consent to participate

All patients included in this study provided written-informed consent to an Institutional Review Board (IRB)-approved prospective biomarker study at the University of North Carolina at Chapel Hill Cancer Hospital that was conducted in accordance with the U.S. Common Rule.

Study design and eligibility

Patients with HPV-associated OPSCC were enrolled on an IRB-approved (UNC IRB 11-1924) prospective biomarker study (NCT0316182). All patients provided written-informed consent. The major eligibility criteria were (i) 18 years age or older, (ii) biopsy-proven squamous cell carcinoma of the oropharynx, (iii) p16 positivity (defined as >70% of carcinoma cells showing nuclear reactivity), (iv) T0 to T4, N0 to N3, M0 (AJCC 7th edition), and (v) receiving definitive CRT. Patients with T0 to T3, N0 to N2c (AJCC 7th edition), and with a tobacco smoking history of ≤ 10 TPY or >10 TPY and not actively smoking at time of diagnosis were coenrolled on two institutional single-arm phase II deintensified CRT trials (NCT02281955 or NCT03077243). Fifty-five healthy volunteers and 60 patients with non-HPV-associated malignancy were also enrolled as a control cohort. A REMARK diagram of the patient cohorts analyzed in this study is provided in Fig. 1.

Definitive CRT

Patients not coenrolled on the phase II deintensification CRT trials received standard-of-care 70 Gy intensity-modulated radiotherapy (IMRT) with concurrent chemotherapy. All patients enrolled on the phase II deintensification CRT trials received 60 Gy IMRT with concurrent weekly intravenous cisplatin 30 mg/m². Patients with T0–T2 N0–1 disease did not receive chemotherapy. All patients had a 3-month posttreatment PET/CT and clinical examination to determine need for neck dissection. Thereafter, patients underwent clinical examinations every 2 to 4 months for years 1 to 2, then every 6 months for years 3 to 5. Chest imaging was performed every 6 months.

Blood collection and extraction of circulating DNA

Deidentified blood specimens were collected before treatment, weekly during CRT, and with each posttreatment follow-up visit in 10 mL cell-free DNA (cfDNA) BCT blood collection tubes (Streck 218962), and double-spun plasma (2,000 × g) was harvested within 3 days for storage at –80°C for ctHPVDNA analysis. Plasma cfDNA was extracted from 2 to 5 mL of plasma using the QIAamp circulating nucleic acid Kit (Qiagen; Catalog No. 55114). The amount of purified cfDNA was quantified with a Qubit fluorometer and PicoGreen quantification reagents (Invitrogen).

Chemicals and reagents

Droplet digital PCR reagents [2x ddPCR Supermix for Probes (No dUTP) Catalog # 186-3024; Pipet tips Catalog # 186-4121; Cartridge Catalog # 186-4109 and Sealing foil Catalog #181-4040] were purchased from Bio-Rad. Eppendorf 96-Well twin.tec PCR Plates (Catalog #E951020362), Falcon 15 mL Conical Centrifuge Tubes (Catalog # 352096), Falcon 50 mL Conical Centrifuge Tubes (Corning #352098), Qubit dsDNA HS Assay

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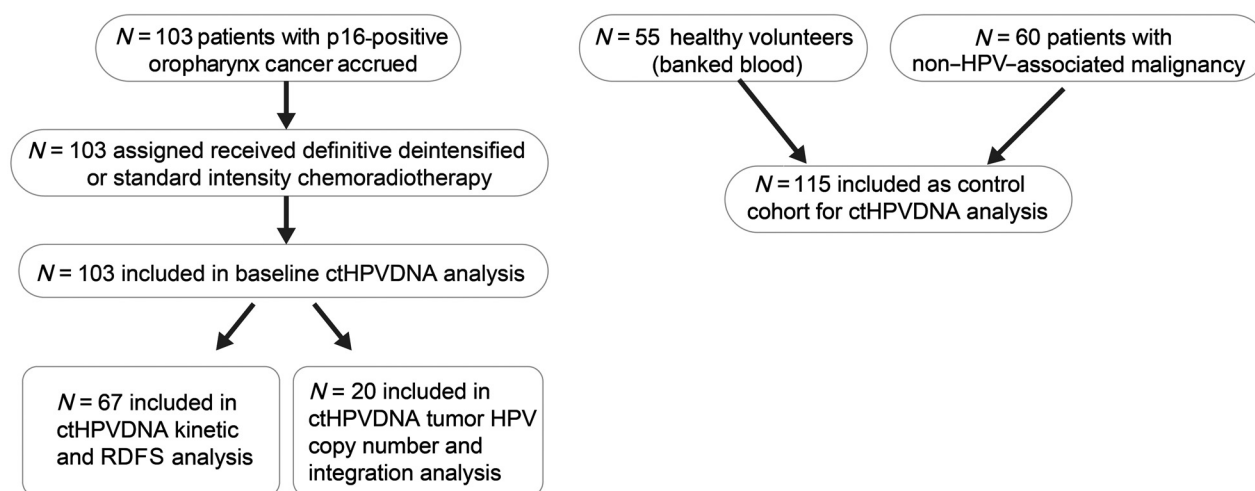


Figure 1.
REMARK diagram of patient cohorts analyzed in this study.

Kit (Catalog # Q32851), and Qubit Assay Tubes (Catalog # Q32856) were purchased from Fisher Scientific. BSA was from Sigma (Catalog # A7906; Sigma–Aldrich). Circulating DNA was quantified with Qubit (Invitrogen). Target gene fragments for HPV subtypes 16, 18, 31, 33, and 35 were synthesized as gBlocks from Integrated DNA Technology (IDT) for use as positive controls (Supplementary Table S1). These gBlocks were cloned into a pCR2.1-TOPO TA vector by Topo TA cloning kit (Invitrogen) according to the manufacturer's protocol. The gene fragment constructs were verified by Sanger sequencing (Genewiz). FAM-, TET-, or HEX-conjugated Locked Nucleic Acids–modified DNA oligonucleotide probes and FAM-ZEN dual quencher probes were synthesized by IDT (see Supplementary Table S1).

ctHPV16DNA dPCR assay and detection of HPV variants

Primers and 5' hydrolysis probes were designed to specifically detect an approximately 75 bp amplicon within the E7 gene encoded by high-risk HPV strains 16, 18, 31, 33, and 35. Sequence details of the primers and probes used in this study are listed in Supplementary Table S1. Each reaction assay contained 10 μ L of 2x dPCR Supermix for Probes (No dUTP), 0.9 μ mol/L of respective primers, 0.25 μ mol/L of respective probes, and 5 to 50 ng of cfDNA in a final volume of 20 μ L. dPCR assays were performed on the QX100 and/or QX200 platforms outfitted with an automated droplet generator (Bio-Rad). Each assay was validated against control DNA samples comprising either the complete genome for the HPV strain of interest, or the relevant portion of the E7 gene. All assays were specific for the HPV strain of interest, with no cross-reactivity to all other HPV strains tested. Standardized thresholds for positive and negative droplets were used across all samples. A dPCR assay for a control locus in the *ESR1* gene was used to assess sample quality. All dPCR data were analyzed using QuantaSoft software version 1.7.4.0917 (Bio-Rad). Additional details of the dPCR assays used in this study are provided in the Supplementary Material.

Analysis of tumor HPV copy number and integration status

Targeted exon sequencing was conducted after IRB approval through the UNCseq (University of North Carolina,

Chapel Hill, NC, http://www2.cccc.unc.edu/~unchreg/UNCseq_NCT01457196) pipeline, as previously described (22, 23). In brief, sequencing data are routed through an automated pipeline. This workflow uses paired tumor and normal libraries to detect somatic mutations, large and small indels, structural variants, and copy-number aberrations. Raw sequences are aligned to hg19_(hs37d5), supplemented with the HPV16 (NC_001526.2) and HPV 18 (NC_001357.1) genomes, using the Burrows–Wheeler Aligner "mem" algorithm. The alignments are then refined using our Assembly Based ReAlignment (ABRA; ref. 24) procedure. ABRA generates a list of structural variants found in target locations, including HPV–human fusions, and permits alignment of both junction-containing and junction-bridging reads. HPV–human chimeric reads were distinguished from non-chimeric HPV alignments, and these counts were normalized to the count of nonchimeric human alignments. Alignment details from ABRA were integrated with these normalized counts to generate circos plots for a subset of samples (25).

Statistical analyses

Linear regression, *t* tests, and survival analyses were performed in Prism 8 (Graphpad) software. Regional disease-free survival (RDFS) was estimated using the Kaplan–Meier method. RDFS was measured from the time of the first dose of CRT and was defined as time to detection/development of persistent/recurrent disease in cervical lymph nodes. Two-sided log-rank test was applied to compare RDFS of different subgroups.

Results

Patient characteristics

Between February 2016 and August 2018, 103 patients were enrolled. The clinical characteristics of the study population are shown in Table 1, and a REMARK diagram of cohorts for study analyses is presented in Fig. 1. A total of 87 patients (84%) received deintensified CRT on clinical trial (60 Gy). The majority of patients were never smokers or had ≤ 10 pack years of tobacco use (75%). HPV status was unknown in 48% (all tumors were p16 positive).

Table 1. Patient characteristics

	N = 103	%
Age (mean)	60	NA
Gender		
Male	92	89%
Female	11	11%
Tobacco use		
Never smoker	56	54%
≤10 pack years	21	21%
>10 pack years	26	25%
T stage		
T0	5	5%
T1	15	14%
T2	69	67%
T3	7	7%
T4	7	7%
N stage		
N0	5	4%
N1	16	16%
N2	82	80%
N3	0	0%
HPV tissue status		
HPV ⁺	44	43%
HPV ⁻	10	9%
HPV unknown	49	48%
Radiation dose		
60 Gy	87	84%
70 Gy	16	16%
Chemotherapy		
Yes	87	84%
No	16	16%

Cancer control outcomes

The median follow-up for 67 patient subset for whom weekly blood samples were collected and a posttreatment PET scan was available was 16.5 months (range, 3.9–32.9). Ten patients had a neck dissection because of incomplete radiographic response on the 3-month posttreatment PET/CT. Five of 10 patients had a positive neck dissection specimen. Three patients had recurrent disease: 0 local, 2 regional and distant, and 1 distant only. The actuarial 1-year local control, regional control, and distant metastasis-free survival, and overall survival were 100%, 91%, 94%, and 98%, respectively. Actuarial 1-year estimate of RDFS (5 positive neck dissection specimens and 2 regional recurrences) was 91%.

Plasma circulating tumor HPV DNA is detectable in a majority of oropharyngeal cancer patients

A dPCR assay was developed to amplify and quantify a 73 bp region of the HPV16 E7 gene. The assay does not cross-detect other HPV subtypes including HPV-6/11/18/31/33/35 (Supplementary Fig. S1A). A titration analysis reveals exceptional linearity ($R^2 = 0.99$) and low variance over 5 orders of magnitude (Supplementary Fig. S1B).

ctHPV16DNA was undetectable in 52 of 55 healthy volunteers (Fig. 2A). Three female volunteers, ages 20, 21, and 31, had low but detectable levels of ctHPV16DNA. We were unable to access the clinical history of these healthy volunteers to establish cervical HPV status. ctHPV16DNA was not detected in plasma DNA extracted from 60 patients with non-HPV-associated malignancy (Fig. 2A). Eighty-four of 103 (82%) patients with HPV-associated OPSCC had detectable pretreatment ctHPV16DNA (Fig. 2B), with median copy number of 419 copies/mL plasma (range, 8–22,579). The 19 samples with undetectable ctHPV16DNA were also analyzed by specific dPCR assays for alternative high-risk

HPV strains (HPV 18/31/33/35; Supplementary Fig. S2). Eight of 19 patients with undetectable baseline ctHPV16DNA were positive for ctHPV DNA from an alternative high-risk HPV strain (one HPV31, three HPV33, and four HPV35), with median copy number of 124 copies/mL (range, 71–15,829). Four of the remaining 11 patients who were negative for ctHPV DNA for all high-risk strains tested were analyzed by tumor genomic sequencing, and 2 were found to have a TP53 mutation. Based on the analysis of 103 cases and 115 controls, we estimate that ctHPV DNA testing using an optimized multianalyte dPCR assay has 97% specificity and 89% sensitivity to identify patients with newly diagnosed and nonmetastatic HPV-associated OPSCC.

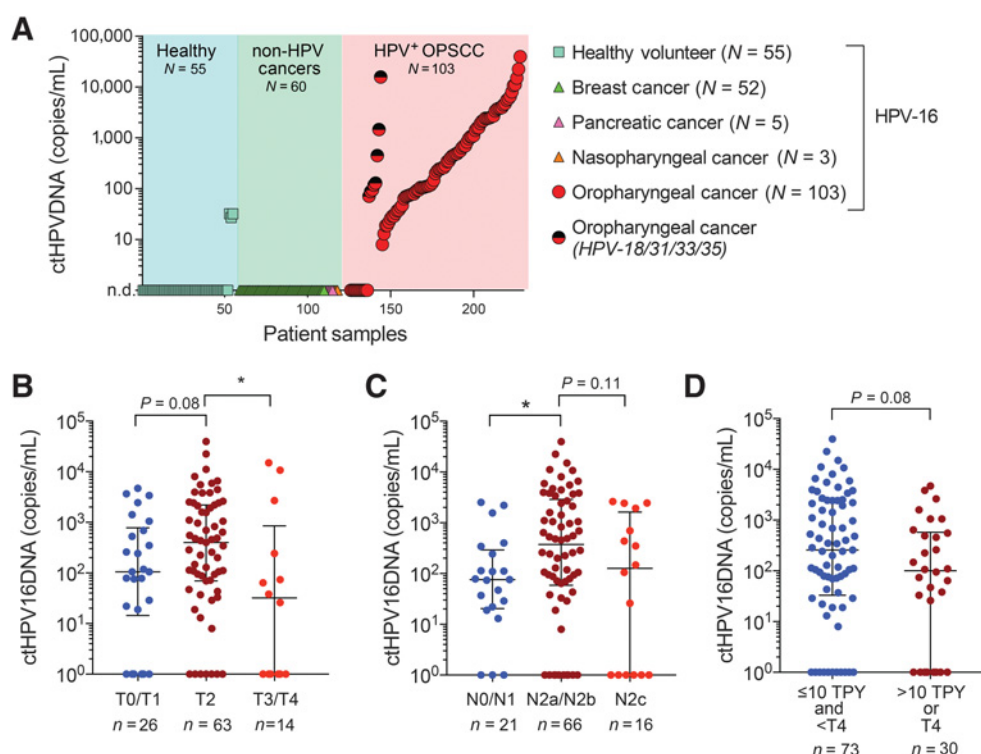
Baseline ctHPV16DNA levels correlate with tumor burden and clinical risk factors

There was a trend toward higher baseline plasma ctHPV16DNA levels in patients with T2 tumors compared with T0/T1 tumors (Fig. 2B). Despite having a relatively low number of T3/T4 tumors in the cohort, these patients had significantly lower baseline levels of ctHPV16DNA relative to patients with T2 tumors (Fig. 2B), suggesting that larger tumor size may be associated with lower rates of ctHPV DNA release. Consistent with a prior study (12), we observed significantly higher ctHPV16DNA levels in patients with N2a/N2b versus N0/N1 disease (AJCC 7th edition; Fig. 2C). Again, there was a trend toward lower baseline ctHPV16DNA in patients presenting with N2c disease, relative to the N2a/N2b patient subgroup (Fig. 2C). Similar trends were seen when AJCC 8th edition N stage classifications were used (Supplementary Fig. S3). These findings suggest that tumor burden alone may not explain the variable levels of pretreatment ctHPV16DNA in OPSCC patients. Notably, patients with adverse clinical risk factors (>10 TPY or T4) had lower baseline ctHPV16DNA levels than patients with low clinical risk (≤10 TPY and <T4; Fig. 2D). Thus, although ctHPV16DNA is detectable in a majority of patients with newly diagnosed HPV-associated OPSCC, low baseline and undetectable ctHPV16DNA levels may be associated with clinically higher-risk disease.

Rapid clearance kinetics of ctHPV16DNA during CRT in a subset of patients

We next investigated kinetics of ctHPV16DNA clearance during CRT in a subset of 67 patients for whom weekly blood samples were collected and a posttreatment PET scan result was available to assess treatment response (median follow-up 16.5 months). Fifty-four of 67 patients had detectable baseline ctHPV16DNA. In 35 of 54 patients (65%), ctHPV16DNA levels increased after initiating CRT before diminishing later in the course of therapy (Fig. 3A). Patients with ≤200 copies/mL of baseline ctHPV16DNA had labile clearance kinetics (Supplementary Fig. S4), possibly due to a lower reliability of signal detection or nonuniform responses to treatment. Thus, we analyzed weekly ctHPV16DNA clearance kinetics in the 40 patient subset (out of 67) who had >200 copies/mL baseline ctHPV16DNA. Levels of ctHPV16DNA diminished during CRT, and approximately 80% of patients had eliminated all ctHPV16DNA by the end of CRT (Fig. 3B). Furthermore, ctHPV16DNA had cleared in 92%, 94%, and 100% of patients by 6 months, 1 year, and 2 years after CRT, respectively. Patients with detectable ctHPV16DNA at the end of week 6 of CRT did not have a higher incidence of residual or recurrent disease (Supplementary Fig. S5). Alternatively, we measured percent clearance of ctHPV16DNA at week 4

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**Figure 2.**

Detection of plasma ctHPVDNA in HPV-associated oropharyngeal cancer patients. **A**, Measurement of ctHPVDNA copies/mL plasma in 55 healthy volunteers, 60 patients with non-HPV-associated cancers, and 103 patients with HPV-associated OPSCC. Two-toned (red-black) circles denote patients who were negative for ctHPV16DNA but positive for ctHPVDNA from an alternative high-risk HPV strain (-18/31/33/35). **B-D**, Baseline levels of ctHPV16DNA stratified by **(B)** tumor stage, **(C)** nodal stage, and **(D)** clinical risk factors. Median and interquartile range is shown for each graph. *, $P < 0.05$; P values based on a two-sided Mann-Whitney test.

relative to pretreatment levels as a potential indicator of CRT sensitivity, as has previously been investigated for EBV-associated nasopharyngeal cancer (26). Using a median cutoff of >95% clearance at week 4 (days 25–31), we identified 19 patients with a ctHPV16DNA rapid clearance profile (Fig. 3C). These patients had a consistent drop in ctHPV16DNA during CRT, and all patients had a clinical complete response to CRT (Fig. 3C–D). In contrast, the remaining 21 patients had delayed clearance kinetics ($\leq 95\%$ clearance at week 4), exhibited more unpredictable ctHPV16DNA profiles (Fig. 3E), and 3 of these patients had persistence/recurrent disease after CRT (2 positive neck dissections, 1 distant metastasis) (Fig. 3C). Thus, ctHPV16DNA rapid clearance kinetics, defined as >95% clearance by week 4, may be a biomarker of CRT sensitivity and greater likelihood of disease control.

Low baseline ctHPV16DNA levels correlate with low tumor HPV copy number and HPV integration

Twenty HPV16-positive patients in our cohort were enrolled on an institutional cancer genomic registry study (NCT01457196) and underwent next-generation sequencing (NGS) of their diagnostic tissue biopsy using the UNCSec platform (27), as well as quantification of HPVDNA using dPCR. We observed excellent correlation between HPV16 copies per haploid genome measured by dPCR and HPV16 read counts per 10^6 total aligned reads by NGS (Fig. 4A). Patients with low baseline ctHPV16DNA (≤ 200 copies/mL) had a significantly lower tumor

HPV copy number than patients with high baseline ctHPV16DNA (>200 copies/mL; Fig. 4B). We next evaluated HPV structure in tumors using a validated insertion–deletion detection algorithm (24) to quantify HPV–chromosomal fusion reads that are indicative of HPV integration into the somatic genome (Fig. 4C). As a positive control, we detected end-to-start fusions of the HPV genome ("HPV-HPV fusion") in all samples, which is expected because the HPV reference genome is deposited as a linear sequence yet originates as a circular episome in nature. There was evidence for HPV integration in 8 of 20 patients (40%, Fig. 4C). Circos plots are shown for two representative tumors without and with HPV integration (Fig. 4D). A significantly higher proportion of patients with low tumor HPV copy number (≤ 5 copies/haploid genome) had HPV integration (Fig. 4E, $P = 0.02$). Similarly, patients with low baseline ctHPV16DNA (≤ 200 copies/mL) were more likely to have HPV integration (Fig. 4E, $P = 0.11$). Thus, low baseline levels of ctHPV16DNA are indicative of lower tumor HPV copy number and a greater likelihood of HPV integration, both of which represent adverse tumor genomic features.

ctHPV16DNA rapid clearance profile and correlation to local and regional disease control

We next analyzed correlation between ctHPV16DNA clearance profile and disease control in the same 67 patient subset with available weekly blood samples, posttreatment PET scans, and subsequent clinical follow-up (median 16.5 months). We defined

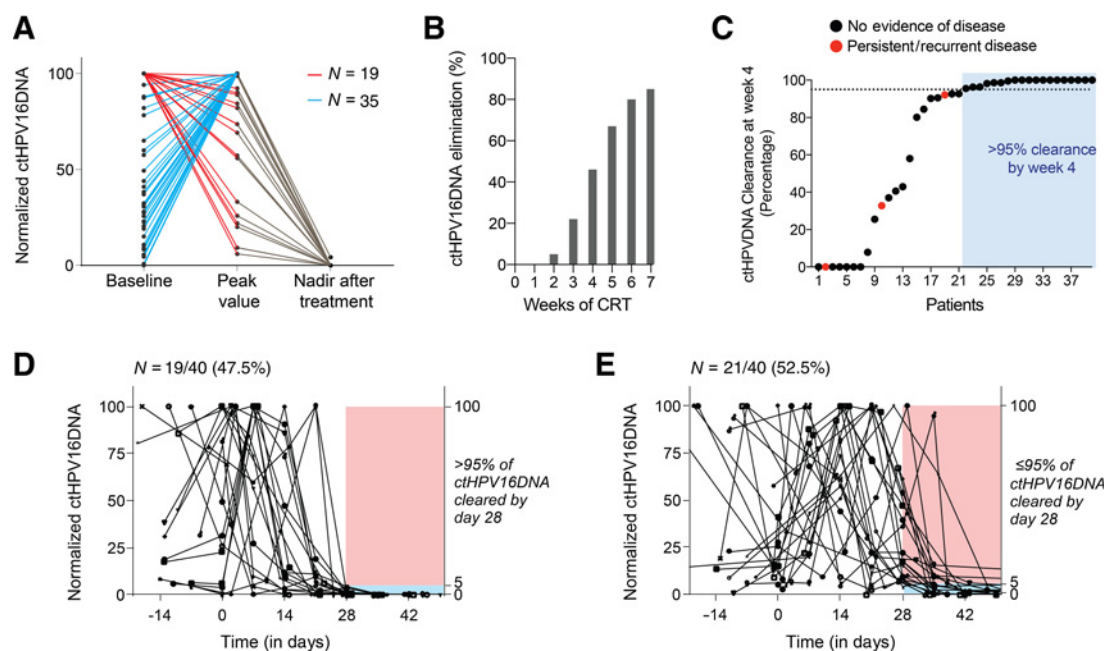


Figure 3.

Variable kinetics of ctHPV16DNA clearance during chemoradiotherapy. **A**, Normalized ctHPV16DNA abundance (relative to the highest measured value for each patient) at baseline, peak value between weeks 1 and 3 of CRT, and nadir after treatment (within 6 months of completing therapy) for 54 evaluable patients with detectable ctHPV16DNA. In one subgroup (red lines, $N = 19$), ctHPV16DNA levels decrease immediately after initiating CRT, whereas in the second subgroup (blue lines, $N = 35$), ctHPV16DNA levels initially spike followed by a subsequent decrease. **B**, Percentage of patients with ctHPV16DNA elimination at different time points after initiating CRT. **C**, Percentage clearance of ctHPV16DNA at week 4 in 40 patients with baseline ctHPV16DNA levels >200 copies/mL. Red dots indicate patients that later developed residual or recurrent disease. **D**, Rapid ctHPV16DNA clearance kinetics in patients with $>95\%$ clearance of baseline ctHPV16DNA levels by week 4 of CRT. **E**, Delayed ctHPV16DNA clearance kinetics in patients with 95% or less clearance of baseline ctHPV16DNA levels by week 4 of CRT.

a favorable ctHPV16DNA clearance profile as high baseline level (>200 copies/mL) with rapid clearance ($>95\%$ clearance by week 4). Twenty-eight percentage of the patient cohort (19/67) had a favorable ctHPV16DNA clearance profile, which did not differ based on established clinical risk factors (Fig. 5A). The remaining patients had an unfavorable ctHPV16DNA profile (undetectable or ≤ 200 copies/mL at baseline, or $\leq 95\%$ clearance by week 4). Patients with an unfavorable ctHPV16DNA clearance profile had a higher frequency of regional disease persistence/recurrence and distant metastasis, particularly among the clinical high-risk patient subset (Fig. 5B). We next quantified actuarial persistent/recurrent RDFS after CRT. Patients with a favorable ctHPV16DNA clearance profile had 100% RDFS (Fig. 5C). Among patients with an unfavorable ctHPV16DNA clearance profile, there was an interaction with established clinical risk factors. Patients with clinical low-risk disease (≤ 10 TPY and $<T4$) had approximately 90% RDFS, whereas patients with clinical high-risk disease (>10 TPY or $T4$) had approximately 65% RDFS at 18 months after CRT (Fig. 5C, $P = 0.0049$). These findings suggest that favorable ctHPV16DNA clearance profile identifies a low-risk subset of HPV-associated OPSCC patients (possibly regardless of clinical risk factors) that are highly responsive to definitive CRT.

Discussion

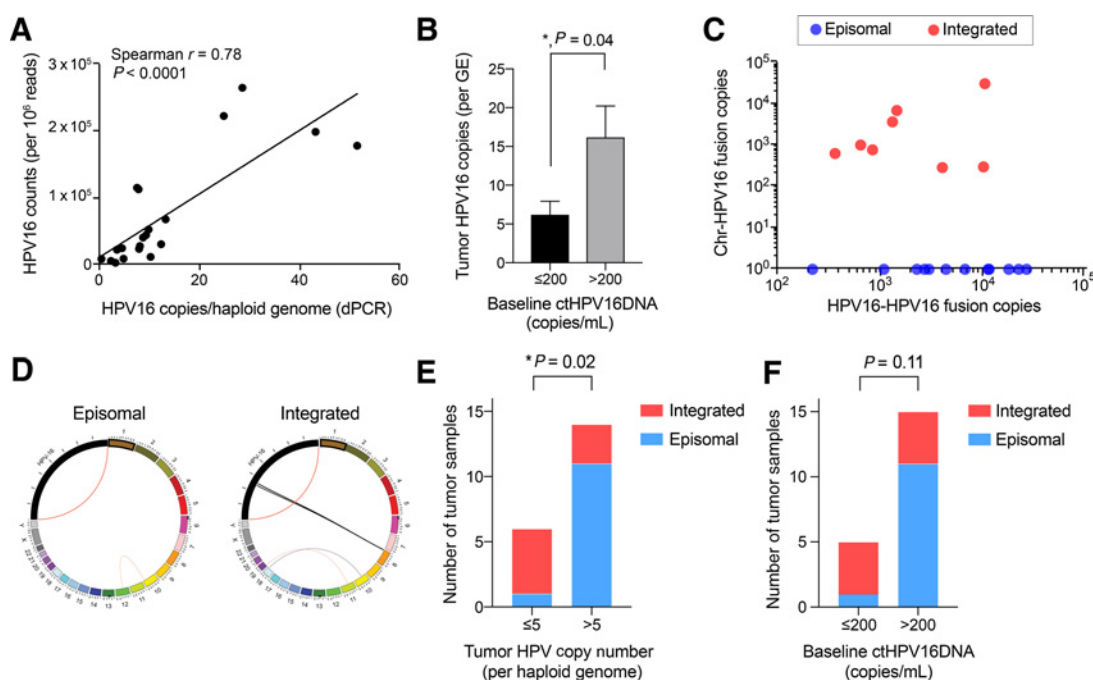
We have prospectively analyzed ctHPV16DNA baseline levels and clearance kinetics in a cohort of newly diagnosed HPV-associated OPSCC patients treated with definitive CRT, 84% of whom were

treated with deintensified CRT on a multi-institutional phase II study. Our findings add to a growing body of evidence that ctHPV16DNA is a promising biomarker for early detection of HPV-associated OPSCC (sensitivity 89% and specificity 97%), similar to the utility of ctEBV16DNA in screening programs for EBV-associated nasopharyngeal cancer (15). Significantly, 11% of our cohort had undetectable ctHPV16DNA from 5 of the most common HPV strains that have been linked to OPSCC, suggesting that some of these cases may represent false positives due to p16-based identification, consistent with another recent report (28).

Although baseline ctHPV16DNA copy number correlated with disease burden, we observed paradoxically lower plasma ctHPV16DNA levels in patients with adverse clinical risk factors (Fig. 2D, >10 TPY or $T4$ disease). By analyzing matched tumor genomics, we discovered that patients with low pretreatment ctHPV16DNA levels (≤ 200 copies/mL) had lower tumor HPV copy number and a higher likelihood of HPV integration, which are associated with worse outcomes in OPSCC (5, 6, 8–12, 29). In contrast, patients with abundant pretreatment ctHPV16DNA levels (>200 copies/mL plasma) were most likely to have tumors with high copy and episomal HPV, which correlates with favorable prognosis in both cervical cancer and OPSCC (8, 9, 11, 14). Thus, beyond simply a measure of tumor burden, high baseline plasma ctHPV16DNA levels (>200 copies/mL) are indicative of favorable tumor genomic biomarkers.

We also found that the rate of ctHPV16DNA clearance correlates with CRT sensitivity. Using week 4 of CRT as a benchmark timepoint, we defined a favorable ctHPV16DNA clearance profile

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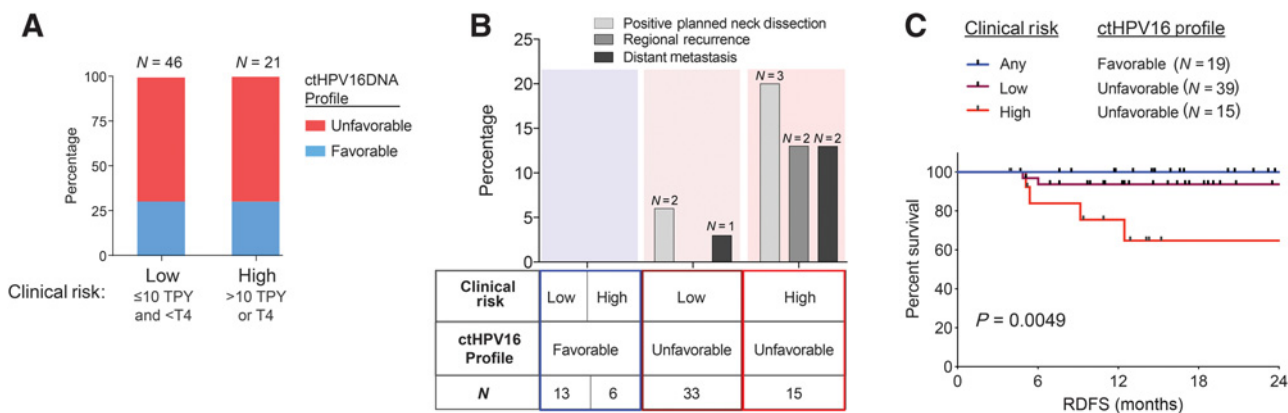
**Figure 4.**

Correlation between ctHPV16DNA and tumor HPV genomic features. **A**, Spearman correlation between copies of HPV16 detected in tumor biopsy samples by dPCR and NGS. **B**, Tumor HPV16 copies per genome equivalent (GE) in patients grouped by baseline ctHPV16DNA levels. Mean \pm SEM is shown. *P* value calculated using a two-tailed, unpaired *t* test with unequal variances. **C**, Normalized NGS counts for hybrid HPV-chromosomal reads (indicative of HPV integration) relative to HPV-HPV fusion reads (positive control). **D**, Circos plots illustrating the detecting HPV and chromosomal rearrangements in a tumor with purely episomal HPV (left plot) and HPV integration into a locus on chromosome 8 (right plot). **E** and **F**, Number of tumor samples with integrated versus episomal HPV after grouping patients by (**E**) tumor HPV copy number (per GE) or (**F**) baseline ctHPV16DNA (copies/mL). *P* values calculated by a two-tailed Fisher exact test.

as elevated baseline levels (>200 copies/mL) that are rapidly cleared (>95% clearance of pretreatment levels by week 4). Patients with a favorable ctHPV16DNA clearance profile had excellent RDFS (~100% at 18 months). Among patients with an unfavorable ctHPV16DNA clearance profile, those with adverse clinical risk factors (>10 TPY or T4) experienced a high rate of residual or recurrent regional nodal disease (~35% at 18 months),

which resembles disease control rates after CRT for HPV-negative OPSCC.

Although there are parallels between ctHPV16DNA in OPSCC and ctEBV16DNA in NPC with regards to early detection, our findings highlight a few clinically significant differences as well. First, whereas high pretreatment levels of ctEBV16DNA are associated with worse prognosis in NPC (16), we find that the opposite is

**Figure 5.**

A favorable ctHPV16DNA clearance profile correlates with disease control in OPSCC patients treated with CRT. **A**, Percentages of clinical low-risk and clinical high-risk patients with a favorable ctHPV16DNA profile, which is defined as baseline ctHPV16DNA >200 copies/mL and >95% clearance by week 4. **B**, Percentage of patients within each subgroup who experience a positive neck dissection, regional recurrence, or distant metastasis. **C**, Kaplan-Meier analysis of RDFS stratified by clinical risk and ctHPV16DNA profile. *P* value calculated using a two-tailed log-rank test for trend.

true for ctHPVDNA in OPSCC (i.e., low pretreatment ctHPVDNA levels are associated with worse clinical outcomes). Second, a recent study demonstrated the prognostic significance of ctEBVDNA at the end of CRT as predictive of NPC recurrence (30). We did not observe a similar correlation with ctHPVDNA, as patients with detectable signal at the end of CRT were not enriched for those that later developed residual or recurrent disease. However, a high ctHPVDNA clearance percentage at week 4 (i.e., >95%) was associated with a lower rate of regional disease persistence/recurrence, similar to what has been shown in EBV-associated NPC (26). A possible explanation for these discrepant findings may be due to the highly variable HPV copy numbers per tumor genome that are observed in HPV-associated OPSCC (see Fig. 4A and B). Tumors with high-copy, episomal HPV may have as many as 50 to 100 copies per cell, which may make it possible to detect small amounts of residual ctHPVDNA even when CRT has effectively treated all of the active disease. Thus, the specific ctDNA profiles associated with clinical outcomes may differ according to the distinct clinical and biological features of EBV-associated NPC and HPV-associated OPSCC, respectively.

The potential clinical utility of ctHPV16DNA monitoring in OPSCC is broad, especially in light of recent evidence that some forms of deintensified therapy are not effective for all patients with HPV-associated OPSCC (31, 32). Future studies should prospectively investigate whether ctHPVDNA kinetics can dynamically guide in "real-time" the intensity of CRT that may be required. Can patients with ≤ 10 TPY and a favorable ctHPV16DNA profile be further deescalated (reduce radiation to 50 or 54 Gy)? Is deintensified CRT efficacious in patients with >10 TPY and a favorable ctHPV16DNA profile? Our data suggest that deintensified CRT should not be offered to patients with an unfavorable ctHPVDNA profile and >10 TPY or T4 disease. Our findings illustrate that dynamic monitoring of ctHPVDNA during CRT may facilitate personalized treatment decisions to maximize tumor control while also minimizing treatment-associated morbidity.

Another potential clinical utility of dynamic ctHPVDNA monitoring may be in the posttreatment, surveillance setting. Our findings indicate that the sensitivity of ctHPVDNA appears to be quite high, and its use to detect cancer recurrence after definitive treatment may be advantageous. ctHPVDNA-based surveillance may detect recurrences earlier than standard clinical visits and occasional radiographic surveillance. Early detection may result in a greater occurrence of the oligometastatic state and thus allow for more aggressive salvage treatment (e.g., metastasectomy, radio-surgery) that may have greater efficacy. Furthermore, ctHPVDNA surveillance may be more cost effective than current surveillance approaches.

Limitations

Although our trial included over 100 patients, it is underpowered due to low occurrence of disease persistence/recurrence and limited follow-up. Despite these limitations, we observed an early trend in worse RDFS (Fig. 5C) in patients with >10 TPY and unfavorable ctHPVDNA kinetics. Validation of our findings using a clinical-grade test in independent and larger patient cohorts will be necessary to confirm our findings.

Conclusions and future directions

Circulating tumor HPVDNA is present in most patients with newly diagnosed HPV-associated OPSCC. Our findings suggest that a rapid clearance profile of ctHPV16DNA clearance during

CRT may have clinical utility in stratifying patients with HPV-associated OPSCC based on likelihood of disease control. Future studies assessing ctHPVDNA as an integral biomarker to guide treatment deintensification are warranted and may facilitate personalized treatment decisions based on tumor biology in addition to clinical risk factors. Finally, prospective evaluation of ctHPVDNA as a biomarker in other HPV-associated malignancies (e.g., cervical and anal cancers) should be evaluated.

Disclosure of Potential Conflicts of Interest

B.S. Chera holds ownership interest (including patents) in and is a consultant/advisory board member for Naveris. S. Kumar is listed as a co-inventor on a patent application regard a method for measuring tumor derived viral nucleic acids in blood samples, which is owned by the University of North Carolina at Chapel Hill and licensed to Naveris. G.P. Gupta holds ownership interest (including patents) in and is a consultant/advisory board member for Naveris. No potential conflicts of interest were disclosed by the other authors.

Disclaimer

The funders had no role in design and conduct of the study; collection, management, analysis, and interpretation of the data; preparation, review, or approval of the article; and decision to submit the article for publication.

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