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Significant association between host transcriptome-derived HPV oncogene E6* influence score and carcinogenic pathways, tumor size, and survival in head and neck cancer

Tingting Qin PhD^{1§} | Lada A. Koneva PhD^{1,2§} | Yidan Liu PhD^{1,3§} |

Yanxiao Zhang PhD^{1,4} | Anna E. Arthur PhD^{5,6} | Katie R. Zarins MPH⁵ |

Thomas E Carey PhD⁷ | Douglas Chepeha MD^{7,8} | Gregory T. Wolf MD⁷ |

Laura S. Rozek PhD⁵ | Maureen A. Sartor PhD¹

Correspondence

Laura S. Rozek, Department of Environmental Health Sciences, University of Michigan, Ann Arbor, MI. Email: rozekl@umich.edu

Maureen A. Sartor, Department of Computational Medicine and Bioinformatics, University of Michigan, Ann Arbor, MI. Email: sartorma@med.umich.edu

Present address

Lada A. Koneva PhD, Kennedy Institute of Rheumatology, University of Oxford, United Kingdom

Yidan Liu PhD, Women's Hospital, School of Medicine, Zhejiang University, Hangzhou, ZhejiangChina

Yanxiao Zhang PhD, Ludwig Institute for Cancer Research, La Jolla, California

Anna E. Arthur PhD, Department of Food Science and Human Nutrition, University of Illinois at Urbana-Champaign, Champaign, Illinois

Abstract

Background: Human papillomavirus (HPV) oncogenes E6, E7, and shorter isoforms of E6 (E6*) are known carcinogenic factors in head and neck squamous cell carcinoma (HNSCC). Little is known regarding E6* functions.

Methods: We analyzed RNA-seq data from 68 HNSCC HPV type 16-positive tumors to determine host genes and pathways associated with E6+E7 expression (E6E7) or the percent of full-length E6 (E6%FL). Influence scores of E6E7 and E6%FL were used to test for associations with clinical variables.

Results: For E6E7, we recapitulated all major known affected pathways and revealed additional pathways. E6%FL was found to affect mitochondrial processes, and E6%FL influence score was significantly associated with overall survival and tumor size.

Conclusions: HPV E6E7 and E6* result in extensive, dose-dependent compensatory effects and dysregulation of key cancer pathways. The switch from E6 to E6* promotes oxidative phosphorylation, larger tumor size, and worse prognosis, potentially serving as a prognostic factor for HPV-positive HNSCC.

KEYWORDS

 $E6,\,E6^*,\,E7,\,head\,\,and\,\,neck\,\,cancer,\,human\,\,papillomavirus,\,influence\,\,score,\,survival$

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¹Department of Computational Medicine and Bioinformatics, University of Michigan, Ann Arbor, Michigan

²Kennedy Institute of Rheumatology, University of Oxford, United Kingdom

³Women's Hospital, School of Medicine, Zhejiang University, Hangzhou, Zhejiang, China

⁴Ludwig Institute for Cancer Research, La Jolla, CA

⁵Department of Environmental Health Sciences, University of Michigan, Ann Arbor, Michigan

⁶Department of Food Science and Human Nutrition, University of Illinois, Urbana-Champaign, IL

⁷Department of Otolaryngology/Head and Neck Surgery, University of Michigan, Ann Arbor, Michigan

⁸Department of Otolaryngology/Head & Neck Surgery, University of Toronto, Toronto, ON

[§]Tingting Qin, Lada A. Koneva, and Yidan Liu contributed equally to this study.

Douglas Chepeha MD, Department of Otolaryngology/Head & Neck Surgery, University of Toronto, Toronto, Ontario, Canada

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1 | INTRODUCTION

Infection with oncogenic human papillomavirus (HPV) is well established as a causative factor for an increasing subset of head and neck squamous cell carcinoma (HNSCC), especially in the oropharynx where HPV may now account for up to 70% of tumors. The HPV "early" proteins E6 and E7 are the main oncogenic proteins leading to HPV-related cancers, repressing immune response and altering the cell cycle program in the host that together allow replication after squamous differentiation and survival despite abnormal mitoses.² HPV E6 and E7 play multiple roles, some of them synergistic, in the tumorigenesis of cancer. E7 is well known to bind to the retinoblastoma tumor suppressor (pRB) and mark it for degradation,3 which results in E2Fdependent gene transcription and further cell cycle (S phase) activation and proliferation.⁴ The loss of pRB activity could lead to p53-dependent apoptosis; however, this is avoided by E6, which induces p53 degradation. E6 recruits the unbiquitin ligase, E6AP, which polyunbiquitinates the p53 tumor suppressor protein for proteosome-mediated degradation, leading to the inhibition of p53-dependent apoptosis and enhanced cell cycle activation.⁵ Besides the p53-dependent apoptosis pathway, E6 and E7 also inhibit alternative apoptosis pathways, including anoikis and the cytokine-activated extrinsic apoptotic pathway.6

E6 and E7 are involved in several pathways in addition to apoptosis and the cell cycle. They cooperate to inhibit the host immune response. It was shown that high-risk HPV E6 interacts with interferon-regulatory factor 3 (IRF-3) and inhibits its activation of downstream factors. Similarly, E7 binds with IRF-1 and blocks its function. Recent studies showed that E6 and E7 could interact with Wnt signaling components and regulate their signaling transduction through the canonical Wnt/β-catenin pathway. Independent of their cooperative functions, E6 and E7 play roles in epigenetic modulation, telomerase activation, DNA damage response, angiogenesis, cell immortalization, and differentiation. $^{11-14}$

E6 has multiple common transcripts, including the longer, full-length transcript, and multiple spliced E6* variants, the most common being a 183 bp exclusion from positions 226 to 409 bp that was only discovered in high-risk HPV subtypes, 15,16 suggesting its role in carcinogenesis. Although the long E6 isoform is more commonly observed to be transcribed from episomal HPV, the shorter E6* variants are more common in tumors with HPV integrated into the host genome, which are the majority of HNSCC cases.¹⁷ E6 splicing is induced by epidermal growth factor depletion, and has been observed to correlate with an increase in TP53 and E7 protein levels and a decrease in pRb. 15,18 Much less is known regarding the function of E6* compared with E6 protein. Evidence suggests that E6* in HPV18 might counteract full-length E6 function, as it could elevate p53 levels in vivo. 19 Recently, E6* was shown to increase reactive oxygen species (ROS) and oxidative stress, which may lead to increased DNA damage.²⁰ In the same study, E6* was observed to decrease superoxide dismutase isoform 2 (SOD2) and glutathione peroxidase (GPX) expression, suggesting a cause for the oxidative stress.

Research on the oncogenic mechanisms of HPV has been performed on a mix of origin cell types (mainly cervical but some oral), HPV types, and stages of HPV infection, and much of it was performed in vitro, sometimes with E6 or E7 transfected in isolation. The extent to which these cumulative findings remain true in oral primary carcinomas infected with HPV16 is not fully known. Although many of the same HPV behaviors and tumor characteristics have been observed in cervical and oropharyngeal HPVassociated cancers, 21,22 it remains unknown whether the findings based on cervical cells can be fully extended to the oropharynx, which has a different tumor microenvironment. Most notably, as opposed to cervical carcinomas, oropharynx tumors arise within or in close proximity to a lymph node and lymphoid follicles. Furthermore, although many direct targets of E6 and E7 are known, the extent that these direct effects extend to downstream pathway events or cause compensatory effects is not known. It is also unclear

whether most effects of E6 and E7 are dose-dependent or act as on-off switches, although it has been shown that E6 degrades procaspase 8 in a dose-dependent manner.²³

Here, we present a genome-wide study of the relationship between combined E6 and E7 mRNA levels, the percent of E6 mRNA that is full-length vs E6*, and host gene expression levels in primary HPV(+) HNSCC cancers. This has allowed us to observe the extensive, downstream effects of the oncogenic HPV proteins on multiple cancer-related pathways. From the set of 18 HPV(+) HNSCC samples collected at the University of Michigan (UM), and 66 HPV(+) HNSCC samples from The Cancer Genome Atlas (TCGA), we selected the total of 68 HPV16 (+) for this study (14 UM and 54 TCGA). Through this analysis, we recaptured all of the main pathways known to be affected by E6 and E7, whether originally described in cervical or oropharynx cells, and identified some novel pathways that are potentially related to the disease pathogenesis. In addition, we propose an influence score to assess the overall impact of E6+E7 (E6E7) and the percent of E6 that is full-length (E6%FL) levels on the host transcriptome, and identified that the E6%FL influence score was inversely significantly associated with tumor mutational burden (TMB) and tumor size, and associated with better overall survival (OS). These associations represent both the direct and indirect effects of E6*, which may involve changes in E7 translation.

2 | MATERIALS AND METHODS

2.1 | Tumor tissue acquisition and sample preparation

HNSCC patients at UM Hospital with untreated oropharynx or oral cavity tumors between 2011 and 2013 were screened for eligibility; those eligible were asked if willing to provide informed consent for collection of tumor tissue and then collected as described previously. 17,24 Tumor tissue and blood were collected into a cryogenic storage tube and flash frozen in liquid nitrogen by surgical staff until storage at -80°C. The flash frozen tissues were embedded in OCT media in vinyl cryomolds on dry ice and stored in −80°C until prepared for histology. H&E slides were sectioned from each frozen tumor specimen on a cryostat and assessed by a board-certified pathologist at the UM for degrees of cellularity and necrosis. Criteria used for inclusion in the study were a minimum of 70% cellularity and less than 10% necrosis; all others were omitted from further study. The first 36 tumors meeting these criteria were selected for RNA sequencing. mRNA library preparation and RNA sequencing were performed as described for these samples in Zhang et al.¹⁷ Briefly, sequencing with an Illumina HiSeq 2000 using 100 nt paired-end reads was performed by the UM DNA Sequencing Core Facility resulting in an average of 47 million reads per sample.

2.2 | UM RNA-seq preprocessing

RNA-seq preprocessing was performed as described in Zhang et al¹⁷ with an exception in the step of differential gene analysis. The raw sequences were aligned to hg19 using Tophat2 v2.0.11²⁵ using default alignment parameters. Quality control was performed using FastQC²⁶ and RSeQC²⁷ before and after alignment. Gene expression levels were quantified using HTSeq v0.6.1p1 with the "intersection-strict" option.²⁸ Normalization and calculation of logCPM values (log counts per million reads mapped) were performed using the *limma* Bioconductor package²⁹ as described below.

The libraries were also aligned to HPV genomes (downloaded from NCBI) using spliced transcripts alignment to a reference (STAR)³⁰ to allow gapped alignment over splicing junctions. STAR first-pass alignment detected the most abundant splice junctions which were then used in the second-pass alignment. All the quantification and analysis below were based on the second-pass alignment. Samples were classified as HPV(+) if they had more than 500 read pairs aligned to any HPV genome, and HPV subtype was determined as the subtype with the most aligned reads. We identified 14 HPV type 16 tumors, 1 type 18, 1 type 33, and 2 type 35. Only the 14 HPV type 16 samples were used in this study. HPV is a small genome with short oncogenes; therefore, in order to optimize the comparability between UM and TCGA data in terms of HPV gene expression and splicing, all UM libraries were trimmed to the first 48 base pairs, which was the read length of all TCGA HNSC RNAseg data. UM RNA-seg data are available at Gene Expression Omnibus #GSE74956.

2.3 | TCGA RNA-seq data reprocessing

RNA-seq fastq files for the 66 TCGA HPV(+) tumor samples were downloaded from CGHub. The data were realigned and analyzed in the same way as UM RNA-seq data described above. The samples contained 55 HPV type 16, 8 HPV type 33, and 3 HPV type 35. One TCGA sample (TCGA-CN-A6V1, Oropharynx, HPV 16 type) was an outlier with extremely low E6 and E7 expression (1.64 CPM), whereas the average expression of E6 and E7 in all other samples was 112.5 CPM. This sample was excluded from downstream analysis due to its outlier expression and to avoid this sample from having a disproportionately high effect on the results. Only the 54 HPV type16 sequencing data were used in this study.

2.4 | Calculating E6E7 expression and E6%FL

The CPM (count per million) values for HPV oncogenes E6 and E7 (heretofore denoted as E6E7 expression) and the percent of full-length E6 (E6%FL) were calculated as previously described. The CPM values were calculated as the number of read pairs aligned to the relevant HPV genome that intersected E6 or E7, divided by the number of million reads in the total library size. E6 alternative splicing results in multiple forms of spliced E6 (referred as E6*). However, the first intron in all forms is missing. To calculate E6%FL, we computed the ratio of average coverage level in the first intron of E6 (approximate full-length level) divided by the average coverage level in the first exon (estimated all E6 mRNA level), referred to as E6%FL. We identified the major donor-acceptor sites of HPV 16 at 227 and 408 bp (for HPV16) by STAR first-pass alignment.

2.5 | Association analysis for E6E7 expression and E6%FL vs host gene expression

The *limma* R Bioconductor package was utilized to examine the association between the host gene expression and E6E7 expression or E6%FL, respectively. Read counts of UM and TCGA mRNA-seq data extracted by HTSeq were transformed to logCPM values by the *limma* function *voom* and analyzed together. Linear regression was applied to calculate the slope and intercept of E6E7 or E6%FL to each host gene expression using the formula:

$$\begin{split} \log \text{ CPM}_g = & \alpha + \beta_0 \times \log \text{ CPM}_{\text{HPV16}} + \sum_{i=1}^n (\beta_i \times \text{covariate}_i) \\ \log \text{ CPM}_{\text{HPV16}} = & \log \text{ CPM}_{\text{E6E7}}, \text{or}, \log \text{ CPM}_{\text{E6\%FL}} \end{split}$$

where g = 1,...,G is a host gene, i = 1,...,n is the covariate, α is the intercept, and β is the slope. P values of the slopes were reported from the limma function lmfit and false discovery rate (FDR) was calculated using the Benjamini-Hochberg method. Genes with FDR < 0.05 were called significant.

2.6 | Backward selection of clinical variables

To identify the optimal set of covariates, we implemented backward selection for high-throughput data.³¹ A total of nine categorical and continuous variables were analyzed: age, smoking status, clinical T classification (the size and

extent of the main tumor, American Joint Committee on Cancer, seventh edition), tumor stage (based on the TNM combination), nodal status, tumor site, sex, HPV integration, and cohort. We grouped certain values of variables that had an insufficient number of tumors. For example, for clinical T classification, we grouped samples as T1 and T2 vs T3 and T4. We analyzed samples located in oropharynx (52 tumors) vs other sites (16 tumors, consisting of 13 oral cavity samples, 1 from larynx, and 2 from hypopharynx). Analysis by tumor stage was performed for groups: stage IV (49 samples) vs other stages (19 samples, consisting of stage I—1 tumor, stage II—10 tumors, and stage III—8 tumors). Nodal status was defined as: N0 and N1 (23 tumors: N0 = 16 and N1 = 6) vs N2 and N3 (45 tumors: N2 = 42 and N3 = 3). The backward selection procedure starts with fitting the weighted linear model of voom-transformed read counts using the lmFit function with the empirical Bayes smoothing implemented by the eBayes function for each of the host genes and includes the effects of primary interest (E6E7 or E6%FL), as well as all nine covariates. For E6%FL, the initial statistical model included covariates for all the same variables, except HPV integration status (8 covariates as the start point) because integration of HPV into the host genome is a likely effect of higher expression of E6*. 32 Indeed, we observed a significant difference in E6%FL by viral integration status (Wilcoxon rank-sum test P value = .007). In each iteration, the least relevant covariate was dropped if the number of its significantly associated genes (FDR < 0.05) less than or equal to 0.1%, or if multiple covariates had zero significant genes, they were dropped in the following order: age, smoking status, T classification, tumor stage, nodal status, site, sex, HPV integration, and cohort; and the resulting reduced model was fit again for all genes. This process was repeated until the least relevant covariate resulted in <0.1% associated genes. We also analyzed the correlation of log transformed E6E7 or E6%FL with clinical variables.

2.7 | Gene set enrichment (GSE) testing

To identify biological pathways and cellular processes affected by E6E7 or E6 %FL levels, GSE testing was performed using RNA-Enrich. GSE testing was performed using GSE testing was followed and Molecular Function) and KEGG Pathways. To filter out the closely related GO terms for reporting purposes, we used the R package $GO.db^{34}$ to determine relationships among significant terms. A GO term was filtered if one or more of its parents, children, or siblings had a higher rank in the list. An R/Bioconductor package Pathview was utilized to visualize the list of driving genes in the KEGG pathway (FDR < 0.05) as reported by the association analysis described above.

2.8 | Influence score calculation and association tests between influence score and tumor features

In order to estimate the overall influence of E6E7 or of E6%FL on the affected host genes, we selected the genes that were significantly correlated with E6E7 or E6%FL, respectively, from the association analysis (FDR < 0.05). For positively associated genes, we ranked the samples by each of those genes' voom-transformed expression values. For the negatively associated genes, we ranked the samples in descending order. To calculate the sample-wise E6E7 or E6%FL influence, we summed the ranks of these genes in each sample, centered the sum by the mean, and then scaled it by its SD across samples. We defined the resulting values as the influence scores. Influence scores were used in place of mRNA levels for testing associations with E6E7 or E6%FL for two reasons: (a) protein levels often have poor correlation with their mRNA levels, and we previously observed that HPV oncogene predicted activity scores correlate better than their mRNA levels with known HPV oncoprotein effects¹⁷ and (b) this is consistent with previous reports showing transcription factor activity scores based on their target genes provide better estimates of the protein's activity level than the mRNA level of the protein.³⁶

We evaluated the association of E6E7 or E6%FL influence scores with tumor features and demographic variables by linear regression. Variables used in this analysis included clinical T classification as continuous (1-3, 4a,b), smoking (never vs smoker), HPV integration (negative vs positive), N classification (N0 or N1 vs N2 or N3), stage (stages I-III vs stage IV), anatomical site (oropharynx vs other), sex, age, and cohort (TCGA vs UM). The associations between influence score and tumor/demographic features were analyzed by analysis of variance, and *P* values from the *F*-test were reported.

2.9 | Calculation of TMB and survival analysis

To evaluate the relationship between E6E7 or E6%FL expression levels or influence scores with cancer biology and clinical features, we focused on the TCGA cohort due to the availability of the mutational and survival data. The somatic mutation data (MAF file) and clinical data of the TCGA HNSC cohort were downloaded from the Genomic Data Commons data portal, and the 54 HPV16(+) patients were extracted. We calculated TMB as the total number of somatic mutations in each patient divided by the total number of megabases that the whole exome sequencing covered (mut

count/Mb). The correlation between TMB and E6E7, E6%FL levels or their influence scores were assessed by the Pearson correlation test. OS of those patients was analyzed for two groups: the high or low levels of E6E7 or E6%FL expression or influence scores using the R packages: *survival*³⁷ and *survminer*. The median score or the optimal score cutoff (searched by the *surv_cutpoint* function in *survminer*) was used to define the two groups. Kaplan-Meier estimates of survival were determined, and a *P* value was calculated using a univariate log-rank test. Cox proportional hazards regression models were used for adjustment of clinical covariate variables (sex, age, clinical stage, tumor site, and smoking status).

3 | RESULTS

3.1 | Overview of cohort and data

We used previously published mRNA sequencing results from 18 HPV(+) tumor samples collected at UM and 66 HPV(+) samples from TCGA.39 Among these 84 samples, 69 were infected with HPV16 while the other 15 samples contained HPV33 (9), HPV35 (5) and HPV18 (1) (Table 1). The two patient populations showed no differences by age, sex, stage, or smoking status; however, the TCGA population did have a slightly lower proportion of oropharynx tumors (P = .06; Table 1). The E6 and E7 proteins of different HPV strains have different behaviors and propensity toward carcinogenesis; consistent with these reports, we found that the expression of E6 and E7 were significantly different among HPV strains in the 83 samples (one TCGA outlier was excluded, see the Methods section; Figure 1A), whereas they were not significantly different between the two cohorts (Figure 1B). In line with the above observations, the percent of E6 transcripts that were of full-length (E6%FL) were also significantly different by HPV strains (Figure 1C), but not different by the two cohorts. (Figure 1D). Therefore, to avoid confounding by HPV type, we restricted our analysis to the HPV16 samples, resulting in a total of 68 combined UM and TCGA samples.

Before studying the association of E6* with host genes, we first studied associations of E6 and E7 with host genes and pathways, many of which are previously known. To identify host genes with E6 and E7 dose-dependent expression levels, we calculated the overall E6E7 log normalized read counts and performed multiple linear regression analysis of these E6E7 expression levels with the mRNA-seq results of the 68 samples. E6 and E7 expression levels were combined, due to their extremely high correlation (r = .977 for HPV16 UM and r = .970 for HPV16 TCGA).

3.2 | E6 and E7 expression correlates with extensive cell cycle, DNA replication and other cancer pathway gene expression in HNSCC

To test for association between host genes and E6E7 expression, we first investigated known clinical and

phenotypic variables that may explain additional heterogeneity in the data. The initial regression model included covariates for patient age, smoking status, clinical T classification, tumor stage, nodal status, sex, anatomical site, HPV integration status, and cohort, and the optimal model with three covariates (cohort, HPV integration, and sex) was selected by a backward selection approach

TABLE 1 Demographic and clinicopathologic characteristics of human papillomavirus (HPV)-positive patients from both University of Michigan (UM) and The Cancer Genome Atlas (TCGA) cohort

Parameter	HPV(+) UM	HPV(+) TCGA	sum_all _HPV	HPV-16 UM tumors	HPV-16 TCGA tumors	sum_HPV-16	UM vs TCGA Fisher's exact test <i>P</i> value
	tumors	tumors					
	18	66	84	14	55	69	
Age at diagnosis							
Median (SD)	58.5 (7.3)	57 (9.2)		58 (7.7)	57 (9.7)		.29
Sex							
Male	17	60	77	13	49	62	1.00
Female	1	6	7	1	6	7	
HPV type							
HPV16	14	55	69	14	55	69	.73
HPV18	1	0	1				
HPV33	1	8	9				
HPV35	2	3	5				
Anatomical site							
Oropharynx	17	47	64	13	40	53	.06
Oral cavity	1	16	17	1	12	13	
Larynx	0	1	1	0	1	1	
Hypopharynx	0	2	2	0	2	2	
Tumor stage							
I	0	2	2	0	1	1	.75
II	1	10	11	1	9	10	
III	2	7	9	2	6	8	
IV	15	47	62	11	39	50	
T stage							
T1	1	8		1	5	6	.73
T2	7	31		6	28	34	
T3	3	10	13	2	9	11	
T4	7	16	23	5	12	17	
N stage	•	-		-	_	•	
N0	1	18	19	1	15	16	.16
N1	2	6	8	2	4	6	.10
N2	11	39	50	10	33	43	
N3	4	2	6	10	2	3	
Smoking status	T		0	1	2	3	
Current	3	13	16	3	10	13	
Former	3 11	30	41	3 7	26		.64
Never	4	22	26	4	26 19	33 23	.04

(Supplementary Table S1).³¹ A total of 261 host genes (228 upregulated and 33 downregulated) were significantly associated with E6E7 expression (Supplementary Table S2; FDR < 0.05). Among the 228 positively associated genes, 40 were involved in DNA replication; 21 genes were involved in cell cycle checkpoint, whose activation are essential for cell cycle progression; and 42 were

involved with DNA repair. The top six positively associated genes by P value were: NGFI-A binding protein 1 (NAB1), cell proliferation regulating inhibitor of protein phosphatase 2A (CIP2A), DNA topoisomerase II alpha (TOP2A), timeless circadian regulator (TIMELESS), DLG associated protein 5 (DLGAP5), and cell division cycle 6 (CDC6). These results suggest that higher E6E7

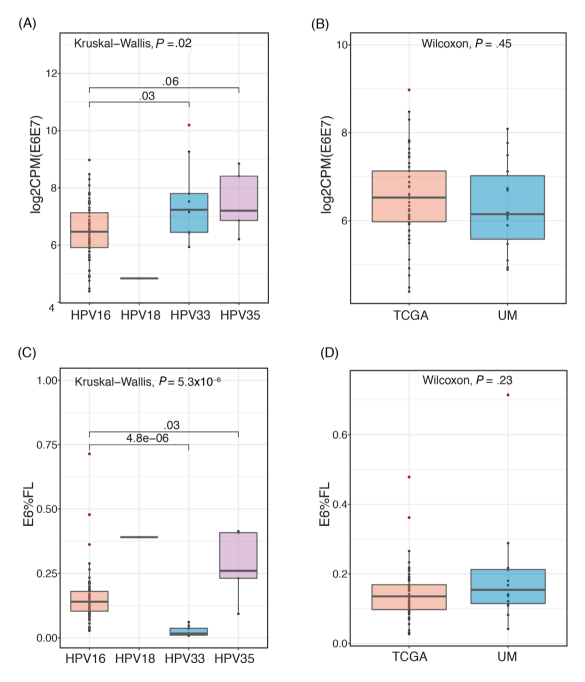


FIGURE 1 A, Box plot showing the normalized RNA-seq expression level of human papillomavirus (HPV) E6+E7, denoted as $\log 2$ CPM (E6E7), in different HPV subtypes across the combined University of Michigan (UM) and The Cancer Genome Atlas (TCGA) 84 HPV-positive head and neck squamous cell carcinoma (HNSCC) samples. B, Box plot showing the expression level of HPV E6E7 in each data cohort (UM and TCGA) (P = .45). C, Box plot showing the normalized RNA-seq expression level of proportion of HPV E6 that is expressed in full-length (E6%FL), in different HPV subtypes across the combined UM and TCGA 84 HPV-positive HNSCC samples. D, Box plot showing the expression level of E6%FL in each data cohort (UM and TCGA) (P = .23) [Color figure can be viewed at wileyonlinelibrary.com]

expression in tumors correlates with enhanced DNA replication and repair, and cell cycle progression, indicating an extensive "dose response" to E6E7 by the host cells.

We identified 94 GO terms and 9 KEGG pathways significantly positively associated, and 83 GO terms and 9 KEGG pathways negatively associated with E6E7

expression (FDR < 0.05; Figure 2A,B; Supplementary Table S3). Consistent with our individual gene results, *cell cycle* and *DNA replication* pathways were found to be enriched with positively regulated genes by E6E7 expression (Figure 2B), and *cell cycle* was the most significant KEGG pathway (odds ratio = 2.33, FDR = 1.04×10^{-21})

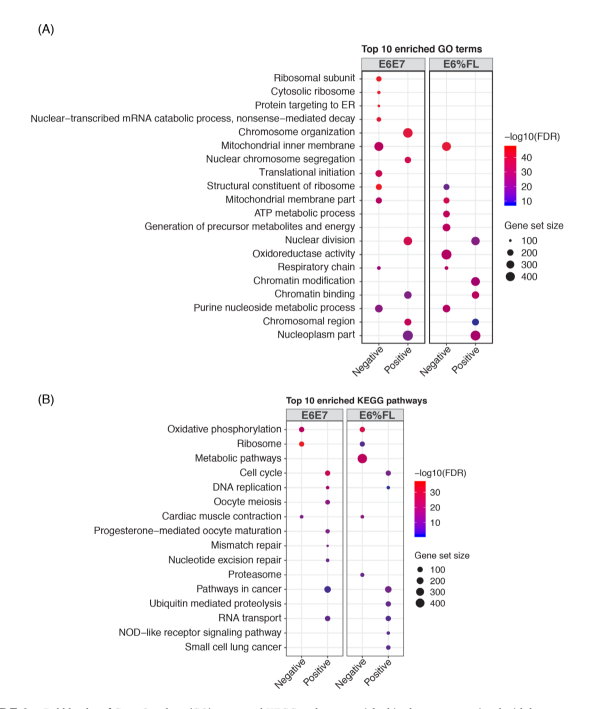


FIGURE 2 Bubble plot of Gene Ontology (GO) terms and KEGG pathways enriched in the genes associated with human papillomavirus (HPV) expression of E6E7 or E6%FL, respectively. A, Ten of the most enriched GOBP (Gene Ontology in biological process domain) terms enriched in host genes positively or negatively associated with E6E7 (or E6%FL) expression. B, Ten of the most enriched KEGG pathways in host genes positively or negatively associated with E6E7 (or E6%FL) expression. The color of the dots denotes the significant levels (reddish: higher significance; bluish: lower significance), and the size denotes the gene set size

with 21 out of 88 genes positively correlated at the FDR < 0.05 level (Supplementary Figure S1; Table S3). Pathways known to be involved in the carcinogenic mechanisms of E6E7 such as regulation of immune response, cell cycle, response to cytokine, Wnt-receptor activity, mismatch repair, nucleotide excision repair, helicase activity, histone binding, and histone kinase activity (Supplementary Table S3) were also found. In addition, novel pathways and GO terms including oxidative phosphorylation (OXPHOS), ribosome, mitochondrial inner membrane, and cardiac muscle contraction were identified as negatively associated with E6E7 expression (Figure 2; Supplementary Table S3). These results confirm the previously reported mechanisms of E6 and E7-related tumorigenesis, 40 and identify novel pathways that may also contribute to E6- and E7-mediated carcinogenesis.

3.3 | Percent of E6 that is full-length as opposed to E6* (E6%FL) is negatively associated with mitochondrial/oxidative processes, ATP metabolic process, keratinization, and inflammation

In high-risk HPV-related carcinogenesis, E6 mRNA is most often spliced to shorter E6 spliced isoforms (E6*), creating truncated proteins whose functions remain elusive. Previous studies suggest E6* can bind to the fulllength E6 and inhibit its function in degrading TP53.41 However, it is known that an increasing percentage of E6* predicts a more severe phenotype in cervical cancer. 42,43 Thus, we hypothesized that E6* has an alternative carcinogenic mechanism that outweighs any tumor suppressive function it may have, such as allowing higher TP53 activity. In order to investigate a possible relationship between E6 splicing and HNSCC carcinogenesis, we determined the percent of E6 expression that was in the form of the full-length isoform for each sample, which is the reverse of percent of E6* (see the Methods section for details). We then performed the same linear regression analysis as above, except using E6%FL instead of E6E7. Similar to the E6E7 analysis, we first investigated known clinical and phenotypic variables that may explain additional heterogeneity in E6%FL (see the Methods section). Using the backward selection process, we identified the optimal model with the covariates of cohort and sex (Supplementary Table S2). A total of 169 differentially expressed genes (156 upregulated and 13 downregulated) were significantly associated with E6%FL from this model (Supplementary Table S4; FDR < 0.05).

GSE analysis revealed 123 GO terms and 20 KEGG pathways negatively associated and 132 GO terms and

20 KEGG pathways (FDR < 0.05) positively associated with E6%FL (Supplementary Table S5). As shown in Figure 2, mitochondrial inner membrane (FDR = 1.2×10^{-43}) and oxidative phosphorylation (FDR = 1.86×10^{-26}) were the top negative terms, with other oxidoreductase-related terms filling the other top negative terms (Supplementary Table S5). These results suggest that mitochondrial/oxidoreductase activity is more activated with a higher rate of spliced E6*, which may lead to heightened oxidative stress and DNA damage. The most highly associated genes related to OXPHOS were NADH:ubiquinone oxidoreductase subunit AB1 (NDUFAB1). However, we did not observe a significant association with either SOD2 or GPX, as was previously observed. Top enriched pathways positively correlated with the E6%FL were chromatin and sequencespecific DNA binding, nuclear division, and microtubule cytoskeleton organization. Also included was stem cell population maintenance, which includes such genes as NOTCH1, SMAD2 & 4, SOX2, STAT3, and FOXO3.

3.4 | Percentage of E6* is positively associated with tumor size and worse survival

Next, we sought to investigate whether the activity level of E6E7 or E6* has any relationship with tumor features or survival. As E6E7 or E6* mRNA expression in each patient may not accurately reflect the protein activity levels, respectively, 17 we calculated an influence score for each tumor sample, defined by the level of influence E6E7 or E6%FL has on the expression of responsive genes. Genes significantly associated with E6E7 expression or E6%FL (FDR < 0.05) were selected to estimate the influence scores (see the Methods section for details). As expected, we observed highly significant correlation between the E6E7 influence score and E6E7 mRNA expression (Figure 3A, left: Pearson's r = 0.68, $P = 2.85 \times 10^{-10}$), as well as between E6%FL influence score and E6%FL expression (Figure 3A, right: Spearman's r = 0.58, $P = 1.68 \times 10^{-7}$). We examined the relationship between E6E7 or E6* and TMB. We hypothesized that as E6* was observed to increase oxidative stress and DNA damage in cells, the percent of E6* would be positively associated with TMB (E6%FL would be negatively associated). We also hypothesized that E6E7 may be negatively associated with TMB, as patients with strong E6 and E7 activity may not require as many mutations for carcinogenesis. To test these hypotheses, we calculated the TMB among the 54 TCGA patients (see the Methods section), and found that TMB had significant negative associations with both influence scores (Figure 3B: E6%FL influence score vs TMB: Pearson's r = -0.36, P = .009; E6E7 influence score vs TMB: Pearson's r = -0.30, P = .03). As

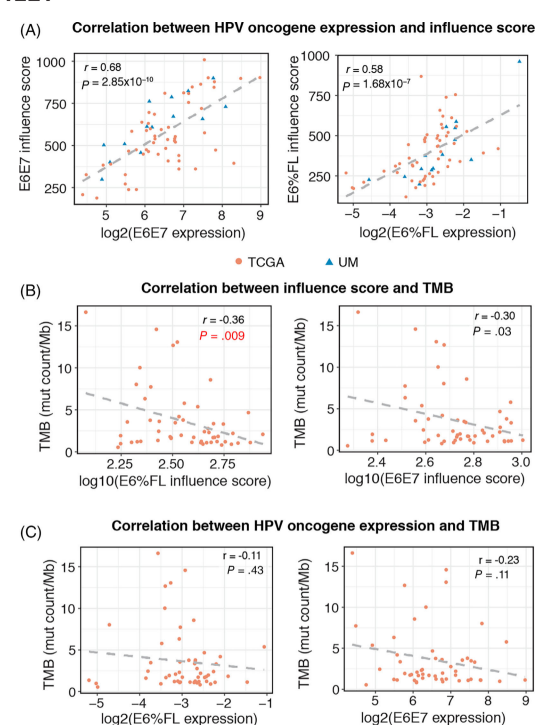


FIGURE 3 A, The correlation between human papillomavirus (HPV) oncogene expression and their corresponding influence scores (left: E6E7 expression level vs E6E7 influence score; right: E6%FL expression level vs E6%FL influence score). The color of dots denotes the cohort (red circle is The Cancer Genome Atlas [TCGA] and blue triangle is University of Michigan [UM] cohort). B, The correlation between the E6E7 or E6%FL influence score and tumor mutational burden (TMB) among the TCGA cohort (left: E6%FL influence score vs TMB; right: E6E7 influence score vs TMB. C, The correlation between E6E7 or E6%FL mRNA expression and TMB among the TCGA cohort (left: E6%FL vs TMB; and right: E6E7 vs TMB) [Color figure can be viewed at wileyonlinelibrary.com]

evidence that the influence scores are more relevant to cancer biology than the mRNA levels, no significant association was observed between TMB and either E6%FL or E6E7 expression levels (Figure 3C).

We then carried out multivariable linear regression analysis to model the association of influence score with different tumor variables, specifically with tumor site (oropharynx [n = 52] vs other [n = 16]); tumor T

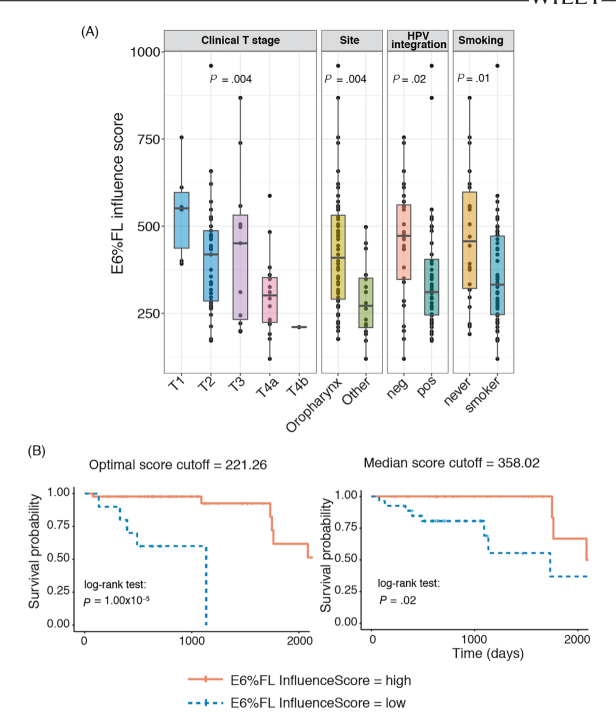


FIGURE 4 The association between E6%FL influence score and tumor characteristics and clinical features. A, Box plot showing the significant difference in E6%FL influence score by clinical T classification (P = .004), tumor anatomical site (P = .003), human papillomavirus (HPV) integration status (P = .005), and smoking status (P = .02). B, The Kaplan-Meier curves showing the significant segregation of overall survival among TCGA patients with HPV16-positive HNSC by E6%FL influence score. Both optimal (left, N = 44 in lower-risk group and N = 10 in higher-risk group at day 0) and median score (right, N = 27 in each subgroup at day 0) cutoffs showed that patients with higher E6%FL influence scores had significantly better survival (Cox proportional hazards: P = .0003 and .02, respectively) [Color figure can be viewed at wileyonlinelibrary.com]

classification (stage I = 6 samples, stage II = 33 samples, stage III = 11 samples, and stage IV = 17 samples; one sample had undefined T classification); HPV integration status; smoking status; sex; tumor N classification; cohort

and age. The E6%FL influence scores demonstrated a significantly negative association with tumor clinical T classification (Figure 4A; Supplementary Table S6; P = .004), meaning that larger tumors were associated with a higher

percent of E6*. Associations resulting in significant unadjusted P values, but which were not significant after multiple-testing adjustment (Figure 4A), included tumor site (P = .004), HPV integration (P = .02) and smoking status (P = .01). On the other hand, the E6E7 influence score had no significant associations after multiple-testing adjustment, and only demonstrated a borderline significant trend with smoking status (P = .04) and tumor site (P = .05).

The significant association between E6%FL influence score and tumor features, especially the negative association with tumor size, triggered us to further assess its clinical relevance. We investigated the OS of the 54 TCGA patients stratified by optimal or median cutoff of E6%FL influence scores (Figure 4B). OS was significantly segregated by both cutoffs (optimal cutoff: log-rank test, $P = 1.00 \times 10^{-5}$; median cutoff: log-rank test, P = .02), and patients with higher E6%FL influence scores, (lower %E6* influence scores), had better survival. In contrast, the mRNA level of E6%FL did not show any association with the patients' survival (Supplementary Figure S2), again suggesting that the E6%FL influence score rather than the E6%FL level itself is more relative to the clinical characteristics. After control for the clinical variables sex, age, tumor stage, tumor site, and smoking status, the significance remained (Cox hazard regression, P = .02 and interquartile range HR = 0.13). In line with the fact that E6* is associated with high-risk HPV(+) cervical cancer, this finding suggests that the E6%FL influence score may also serve as a clinically actionable metric in HNSCC to distinguish patient subtypes, and help guide precision medicine.

4 | DISCUSSION

In this study, we investigated the relationship between HPV oncogene expression and host mRNA expression in HNSCC samples from two cohorts, and used the resulting host gene responses to develop an influence score. This facilitated the identification of the relationship between E6 splicing and tumor size, TMB, and OS. We also confirmed genes and pathways previously identified to be affected by HPV16 E6 and E7, including genes involved in DNA replication, DNA repair and cell cycle, such as CDK2 and CLSPN (claspin), showing that these responses to E6 and E7 expression display a correlative "dose response." HPV16 E7 is known to interact with two cyclin/CDK2 complexes and to inhibit the cyclin dependent kinase inhibitors p21 and p27, leading to the activation of CDK2 and disruption of G1/S cell cycle checkpoint.44,45 CLSPN is essential for DNA-replication stress response, and its degradation is associated with DNA damage checkpoint recovery.46 Another top positively associated gene was *RBL1*, which may share some functional redundancy with pRB based on the high level of sequence similarity between them, suggesting that pRB degradation may activate a compensatory reaction. In addition to the previously known E6 and E7 transcriptional effects, novel genes and pathways were identified, such as mitochondrial function related biological processes. Notably, some Fanconi anemia (FA) pathway genes were found to be significantly upregulated with the increase in E6 and E7 expression, including FA complementation group B (*FANCB*), FA complementation group C (*FANCC*), and FA complementation group M (*FANCM*). This is consistent with the previous finding that patients with FA have much higher risk of HNSCC, ⁴⁷ and HPV(+) HNSCC patients tend to carry more mutations in FA genes. ⁴⁸

E6 mRNA splicing is a critical step in HPV-induced tumorigenesis. However, the function and effect of truncated E6 (E6*) in the disease remains unclear. Genes that significantly correlate with E6 splicing could provide hints of pathways that E6* affects. We did not see any evidence of downstream effects of E6* inhibiting the degradation of TP53, suggesting this may be a weak effect, does not occur in HNSCCs, or does not affect the expression of downstream genes. The negative correlation observed with E6% FL (ie, positive correlation with % E6*), which included OXPHOS, ATP metabolism, mitochondrial membranes, epithelial differentiation (keratinization), and endoplasmic reticulum, suggests that patients with more E6* rely heavily on OXPHOS for energy production. Interestingly, the glycolysis/gluconeogenesis pathway was also found to be negatively associated with percent of E6%FL, although at a lower significance level (FDR = 0.014). This finding appears to contradict the Warburg effect, the tendency of tumors to increase their use of glycolysis for energy production while inhibiting OXPHOS in order to continue growth even in hypoxic conditions. However, an increasing amount of evidence has suggested that OXPHOS is upregulated in some cancers, 49 and one cause of the increased OXPHOS may be the increased mtDNA content in those cancers, including head and neck, esophageal, thyroid, ovarian, prostate, and colorectal cancers.⁵⁰ In addition, the cell differentiation process may induce OXPHOS and mitochondrial biogenesis, increasing ROS production, 51,52 and thus inducing oxidative stress and DNA damage. 20 Taken together, we hypothesize that HPV16 E6* promotes cancer progression and epithelial differentiation via activating the OXPHOS pathway. The findings also suggest that OXPHOS inhibitors may be an effective treatment for E6*-associated high-risk HNSCC patients. Indeed, some recent studies have highlighted mitochondrial metabolism as a target for anticancer therapy.⁴⁹

The higher levels of oxidative phosphorylation and mitochondrial genes may explain the larger tumor sizes at diagnosis among tumors that express a higher percent of E6*. By far, the strongest correlations of host genes with E6* involved mitochondrial functions, including *PDK3*, *FH*, and numerous subunits of ATP synthase, NADH:ubiquinone oxidoreductase, and ATPase H+ transporters. We also showed that the percent of E6* is positively correlated with TMB, suggesting higher E6* may promote carcinogenesis and tumor growth by increasing mutagenesis and/or allowing faster growth via increased energy production. On the other hand, E6* is not associated with E7 at the mRNA level, consistent with the known report that E6* influences translation, not RNA levels of E7.¹⁸ This could help to explain the carcinogenic potential of E6*.

Finally, we found that the percent of E6* is positively associated with worse OS. This finding was uncovered using the E6%FL influence scores, and survival analysis adjusting for multiple covariates. Our findings with E6%FL influence scores, which were not found with E6%FL mRNA expression levels, validate the clinical relevance and potential use of our influence score as a prognostic factor. Hong et al saw a trend in survival based on the ratio of two different shorter isoforms of E6*, E6*I/E6*II. Our definition of E6* included both of these isoforms, but with E6*I being by far the more prominent one.

Overall, this study demonstrates the dose response effects of E6 and E7 oncogenes on the host transcriptome. This suggests that HPV oncogene expression levels are an important indicator of patient prognosis in HNSCC, consistent with findings in cervical cancer.⁵⁴ These analyses identified new genes and pathways affected by E6E7 or the splicing ratio of E6, the latter of which has not been well described. The findings based on the splicing ratio of E6 can guide future studies of the molecular mechanisms underlying E6*-associated carcinogenesis such as mitochondrial metabolism (OXPHOS). The fact that the higher E6%FL influence score was significantly associated with a better OS in HPV16-positive patients further supports E6* being associated with high-risk HPV(+) tumors; we are thus optimistic to propose it as a potential prognostic factor for HPV(+) HNSCC patients although a larger cohort is needed for further validation. One of the limitations of this analysis is that we cannot distinguish between the direct influence of E6*, and the indirect influence through increased translation of E7. Further studies are required to investigate the correlation between E6* and E7 at protein level and explore the underlying oncogenic mechanisms of E6*.

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CONFLICT OF INTEREST

The eligible HNSCC patients at University of Michigan Hospital consented to collect tumor tissue. All authors read and approved of the manuscript. The authors declare no competing financial or nonfinancial interests.

AUTHOR CONTRIBUTIONS

T.Q., L.A.K., and Y.L. performed the bioinformatics and statistical analyses and contributed to the data interpretation of the data and manuscript preparation. Y.Z. contributed specific bioinformatics analyses. A.E.A. and K.R.Z. collected and prepared tumor samples for sequencing, as well as collected clinical data for the UM cohort. D.C. performed the head and neck cancer surgeries and froze the samples. T.E.C. and G.T.W. provided biological inference and clinical interpretation for the results. L.S.R. contributed in sample collection, study design, result interpretation and the manuscript review. M.A.S. supervised the study, determined the bioinformatics and statistical analyses, and participated in the interpretation of data and writing of the manuscript.

ORCID

Tingting Qin https://orcid.org/0000-0003-3810-7578

Thomas E Carey https://orcid.org/0000-0002-5202-7518

Laura S. Rozek https://orcid.org/0000-0001-6731-7000

Maureen A. Sartor https://orcid.org/0000-0001-6155-5702

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of this article.

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