

Patient Burden with Current Surveillance Paradigm and Factors Associated with Interest in Altered Surveillance for Early Stage HPV-Related Oropharyngeal Cancer

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Key Words. Surveillance • Oropharyngeal cancer • Human papillomavirus • Survivorship

ABSTRACT

Introduction. Optimal surveillance paradigms for survivors of early stage human papillomavirus (HPV)-related oropharyngeal cancer are not well defined. This study aimed to characterize patient interest in and factors associated with an altered surveillance paradigm.

Materials and Methods. We surveyed patients with Stage I or II HPV-related oropharyngeal cancer treated at a tertiary care institution from 2016 to 2019. Primary outcomes were descriptive assessment of patient knowledge, interest in altered surveillance, burdens of in-person appointments, and priorities for surveillance visits. Ordinal regression was used to identify correlates of interest in altered surveillance.

Results. Sixty-seven patients completed surveys from February to April 2020 at a median of 21 months since completing definitive treatment. A majority (61%) of patients were interested in a surveillance approach that decreased in-person clinic visits. Patients who self-identified as

medical maximizers, had higher worry of cancer recurrence, or were in long-term relationships were less likely to be interested. Patients reported significant burdens associated with surveillance visits, including driving distance, time off work, and nonmedical costs. Patients were most concerned with discussing cancer recurrence (76%), physical quality of life (70%), mortality (61%), and mental quality of life (52%) with their providers at follow-up visits.

Conclusion. Patients with early stage HPV-related oropharyngeal cancers are interested in altered surveillance approaches, experience significant burdens related to surveillance visits, and have concerns that are not well addressed with current surveillance approaches, including physical and mental quality of life. Optimized surveillance approaches should incorporate patient priorities and minimize associated burdens. *The Oncologist* 2021;26:676–684

Implications for Practice: The number of patients with HPV-related oropharyngeal cancers is increasing, and numerous clinical trials are investigating novel approaches to treating these good-prognosis patients. There has been limited work assessing optimal surveillance paradigms in these patients. Patients experience significant appointment-related burdens and have concerns such as physical and mental quality of life. Additionally, patients with early stage HPV-related oropharyngeal cancers express interest in altered surveillance approaches that decrease in-person clinic visits. Optimization of surveillance paradigms to promote broader survivorship care in clinical practice is needed.

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INTRODUCTION

Early stage human papillomavirus (HPV)-related oropharyngeal cancer represents a distinct entity among head and neck malignancies given better outcomes [1] and differing demographics [2] compared with HPV-negative head and neck cancers. Five-year overall survival rates are in excess of 85% in patients with HPV-related cancers [3, 4], whereas patients with HPV-negative cancers have a 5-year overall survival of less than 50% [1]. In HPV-related cancers, stage I–III patients with a negative positron emission tomography scan at 3 months post-treatment have a 5-year disease-free survival of 91% and 5-year overall survival of 89% [5].

Much contemporary clinical research in these patients currently focuses on treatment deintensification strategies, including alterations in systemic therapies [6] and radiation approaches [7–9], aiming to reduce the known long-term toxicities of treatment (80% peripheral neuropathy [10], 40% ototoxicity [11], 30% nephrotoxicity [12], 15%–20% late dysphagia [13, 14], 15% late xerostomia [15, 16]) and their subsequent impact of quality of life [17, 18]. HPV-related oropharynx cancer-specific surveillance after standard therapy, however, has not been well studied. Current surveillance paradigms offer the same surveillance schedule for all squamous cell carcinomas of the head and neck regardless of HPV status [19], ignoring vastly different outcomes, recurrence patterns, demographics, and comorbidities [20]. Recognizing the lower rates of recurrences in HPV-related cancers as above, revising current surveillance guidelines to deintensify surveillance in good-prognosis patients warrants further evaluation.

Observational epidemiology studies in patients with HPV-related oropharynx cancer suggest that recurrences are frequently detected in the setting of new symptoms, rather than during surveillance visits. Moreover, adherence to standard surveillance paradigms offers limited tangible benefits [21]. Additionally, current surveillance options do not offer a means to detect distant recurrences, which is proportionally more common in this patient population [22].

As we seek ways to improve surveillance options for HPV-related oropharyngeal cancer, patient input is critical. Patient burdens related to appointments and patient desires to address specific concerns during surveillance may illuminate methods to optimize broader survivorship care in addition to standard cancer recurrence surveillance. We aimed to assess patient interest in and factors associated with alternative surveillance approaches, including patient knowledge, burdens of in-person appointments, and priorities for follow-up visits.

MATERIALS AND METHODS

Study Population and Data Collection

Patients who completed treatment for American Joint Commission on Cancer (AJCC) 8 stage I or II HPV-related (assessed by p16 positivity [23]) oropharyngeal cancers without evidence of cancer recurrence and who had been seen at the University of Michigan within the past 6 months

were eligible. All patients were followed with standard surveillance recommendations, which include clinic visits and nasopharyngolaryngoscopy every 3 months for the first 2 years after treatment and every 4–6 months years 3–5. The survey instrument was approved by University of Michigan's Institutional Review Board as part of a larger study following head and neck cancer patients (HUM00042189). Patients were enrolled and consented either while in clinic for a scheduled surveillance visit or via phone and completed the survey electronically. Study data were collected and managed using research electronic data capture tools [24, 25].

Measures

Study participants completed a 38-item survey (see supplemental online Dataset 1). The instrument was based on Andersen's health service utilization model [26], developed using standardized approaches to questionnaire design [27], and based on systematic review of the literature, prior research in patients with head and neck cancer, input from a multidisciplinary team, and survey design experts. Information on patient sex, age, education, race/ethnicity, employment, and income was obtained from the survey. Details on cancer staging and treatment received were obtained from the medical record.

Questions regarding HPV etiology [28], cancer worry [29], self-assessment of health status [30], trust in health care providers [31], shared decision-making preferences [32], health literacy [33], and medical maximizer/minimizer preferences [34, 35] were adapted from previous reports in the literature. Medical minimizer/maximizer preferences distinguish patients who tend to prefer aggressive versus more passive approaches to health care [36]. Concerns related to treatment were adapted from the literature [37] and consisted of 11 topics rated on a three-point scale (not at all, somewhat, and very much); these were used as proxies of items to be addressed during surveillance visits. Surveillance-related burdens were assessed through self-report of time allotted for surveillance visits, method for taking time off work, method of and difficulty of travel to appointments, and money spent on copays and other associated costs (including food, gas, lodging).

The survey presented scenarios to address options for altered surveillance patterns that included decreased clinic visits and assessed interest in nonclinic methods of surveillance. In scenario 1, patients were offered a vignette and asked to rate interest in returning to clinic for fewer routine surveillance visits, from every 3 months in the first 2 years after treatment to every 6–12 months; in scenario 2, patients were then offered a similar vignette with additional education on the expected low risk of recurrence, and again asked to rate interest in altered surveillance. For exploratory analyses, responses for scenario 2 were used as this scenario represents discussions that occur in routine clinical practice. Patients were also asked about potential adjuncts to standard surveillance including blood samples, urine samples, electronic symptom surveys, or expedited symptom-directed survivorship visits (for example, speech-language pathology).

Statistical Analysis

The primary aim of this analysis was to descriptively assess interest in altered surveillance, patient burdens of appointments, and patient priorities for follow-up. Interest in altered surveillance was defined as responses of 4 or 5 on a Likert-type scale from 0 to 5, where 0 represented “not at all interested” and 5 represented “definitely interested.” Wilcoxon signed-rank test was used to compare means between the two scenarios. Treatment-related concerns were analyzed as binary, with “not at all” compared with “somewhat” and “very much.” Exploratory analyses included associating five prespecified variables to avoid overfitting the data (relationship status, worry of cancer recurrence, self-perception of physical health, shared decision-making preferences, minimizer-maximizer preferences) with interest in altered surveillance using ordinal regression. The χ^2 test was used to assess changes in concern with median follow-up time. The data were analyzed using SPSS, version 26 (SPSS, Chicago, IL). Two-sided p values ≤ 0.05 were considered statistically significant.

RESULTS

Sample

Of 90 patients invited to participate, 67 completed surveys, for a 74.4% response rate. Patients received treatment from October 2016 to December 2019, and surveys were completed February to April 2020 at a median of 21.2 months since completing treatment (range, 2.8 to 41 months). Patients had 71.6% AJCC 8 stage I disease, and 73.1% underwent definitive chemoradiation for treatment. Patients had a median age of 60 years (range, 41–83), 92.5% were male, 97% of patients were non-Hispanic White, 53.7% had a bachelor's degree or higher, and 86.6% were in a long-term relationship. Most patients were working at time of survey (61.1%), and nearly half made $\geq \$100,000$ per year (Table 1).

Patients exhibited high levels of self-assessment of physical health, with 64.2% assessing health as excellent or very good. Most patients exhibited preferences for a spectrum of shared decision-making, with only 3% preferring to leave decision-making up to the physician and no patients wanting to make the decision themselves. All exhibited trust in their health care provider (100% yes or definitely yes to some extent). Most patients self-identified as medical maximizers (71.6%), most had low levels of cancer worry (mean, 5.48; SD, 3.56 on 20-point scale), and most reported high levels of health literacy (86.6%, Table 2).

Knowledge

Almost all patients knew that HPV caused their cancer (94.0%), and most agreed that their particular cancer was unlikely to recur (74.6%). Most patients characterized risk of recurrence in either a local or distant location as low or very low (70.1%). Slightly more patients expected a recurrence to be local, although the proportion was near 50% in both: 58% of patients agreed with the statement “if my cancer comes back, it is likely to come back in my throat”; 53% of patients agreed with the statement “if my cancer

comes back, it is likely to come back elsewhere in my body.”

Altered Surveillance

Patients were asked to report interest in decreasing in-person post-treatment surveillance clinic visits. In scenario 1, asking about patient interest in decreased in-clinic surveillance visits, 55.2% of patients were interested in altered surveillance (rated 4–5 on Likert scale), and only 8.9% were not interested at all (rated 0–1 on Likert scale). When offered additional information regarding low risk of recurrence before asking again about interest in decreased in-clinic surveillance visits in scenario 2, 61.2% were interested in altered surveillance (rated 4–5 on Likert scale), and only 7.4% were not interested at all (rated 0–1 on Likert scale). Comparison of responses to both scenarios showed there was no significant difference in responses to scenario 1 (mean, 3.46; SD, 1.50) versus scenario 2 (mean, 3.58; SD, 1.36; $p = .203$).

An exploratory analysis of factors associated with interest in altered surveillance were assessed with an ordinal regression model incorporating prespecified variables of relationship status, worry of cancer recurrence, self-perception of physical health, shared decision-making preferences, and minimizer-maximizer preferences. On multivariable regression, being a medical maximizer (higher on the minimizer-maximizer scale; odds ratio [OR], 0.64; 95% confidence interval [CI], 0.45–0.89, $p = .008$), being in a long-term relationship (OR, 0.12; 95% CI, 0.03–0.56; $p = .007$), and having higher worry of cancer recurrence (OR, 0.86; 95% CI, 0.75–0.99; $p = .041$) were all associated with decreased interest in altered surveillance, whereas physical health ($p = .36$) and shared decision-making ($p = .60$) were not associated with interest in altered surveillance. Although time from end of treatment was not prespecified as a variable of interest, in a separate model, time from the end of treatment was assessed and was not significantly associated with interest in altered surveillance.

Patients were asked to select from four nonclinic based surveillance options as a means to supplement surveillance. When asked to select only one option, 61% selected blood samples as the preferred nonclinic way to follow cancer, 19% selected surveys, 10% selected symptom-directed survivorship visits, and 9% selected urine samples. When allowed to select multiple options as a nonclinic based surveillance option, 94% selected blood samples, 63% selected urine samples, 58% selected surveys, and 48% selected symptom-management visits.

Surveillance Burden

Patients were asked to assess varying burdens related to surveillance appointments. Most patients felt it was easy to get to follow-up appointments, with 59.7% stating “very easy” and 32.8% stating “somewhat easy.” A minority of patients (9.0%) felt that it was somewhat difficult to come to appointments. Patients drove a median of 57 miles to reach appointments (range, 2–346), and 22.4% of patients drove more than 100 miles to come to appointments.

Patients allotted a significant portion of their day to be able to attend surveillance visits. More than 80% of patients

Table 1. Sample characteristics

Characteristics	n (%)
Patient characteristics	
Age, yr	
Median	60
Min	41
Max	83
Gender	
Male	62 (92.5)
Female	5 (7.5)
Smoking status (at time of treatment)	
Never smoker	40 (59.7)
Former smoker	23 (34.3)
Current smoker	4 (6.0)
Race/ethnicity	
Non-Hispanic White	65 (97.0)
Black	1 (1.5)
Prefer not to answer	1 (1.5)
Tumor and treatment characteristics	
AJCC 8 group stage	
I	48 (71.6)
II	19 (28.4)
Primary therapy	
Chemoradiation	49 (73.1)
Radiation	5 (7.5)
Surgery	13 (19.4)
Median time since end of treatment, mo	21.2
Min	2.8
Max	41.0
Socioeconomic characteristics	
Education	
High school or less	8 (11.9)
Some college or trade school	23 (34.3)
Bachelor's degree or higher	53.7)
In long-term relationship	
Yes	58 (86.6)
No	9 (13.4)
Employment status	
Working full-time	34 (50.7)
Working part-time	7 (10.4)
Not working	26 (38.8)
Health insurance	
Yes	66 (98.5)
No	1 (1.5)
Financial dependents	
0–1	46 (68.7)
2+	21 (31.3)
Household income	
Less than \$50,000	9 (13.4)
\$50,000–\$99,999	16 (23.9)
\$100,000 or more	30 (44.8)
I prefer not to answer	12 (17.9)

Abbreviation: AJCC, American Joint Commission on Cancer.

allotted a half day or more to attend a single surveillance visit, with nearly half of patients allotting at least a half day (49.3%) and 26.9% allotting a full working day for one appointment. A minority of patients, 6.0%, allotted more than one full day (this presumably includes an overnight stay for those who drove from long distances). To obtain time off from work to attend these visits, 19.4% used unpaid time off, and 16.4% used sick days, vacation days, or the Family and Medical Leave Act; 41.8% of patients did not work.

Almost all patients, 98.5%, had health insurance. Patients were asked to estimate direct cost to them associated with surveillance visits. Patients were first asked about copays, or immediate out-of-pocket costs associated with the visit itself. Nearly half of patients (46.3%) were unable to identify an amount. In free text, 21 patients stated that they did not know the amount; 2 patients stated that it varies; 1 person gave a percent based on insurance; and 2 patients stated “up to” a certain dollar amount (\$1,000 and \$3,500 for these 2 patients, presumably reflecting insurance deductible amounts). Two patients explicitly stated that they had a very high deductible plan; 19.4% of patients did not pay any copay. The remaining 34.3% reported a range of costs, with 17.9% reporting \$26–\$50 and 7.5% reporting >\$50.

Patients were also asked to estimate out-of-pocket costs related to gas, food, or lodging for each visit; 17.9% of patients were unsure or reported no costs. The remaining 82.1% reported some cost, with 40.3% reporting \$1–\$25 in cost, 19.4% reporting \$26–\$50, and 22.4% reporting >\$50 (Table 3). For all patients who reported a dollar amount for copay or nonmedical out-of-pocket cost, 22.4% of patients reported spending more than \$100. Of note, the costs assessed here did not include medical bills or additional subsequent costs from the health care system or insurers.

Priorities for Surveillance

Treatment-related concerns were elicited from each patient to allow for outlining concerns that could be addressed during surveillance care. Cancer recurrence remained the most important concern, with 76% of patients noting this as somewhat or very important. Mortality was important to 61% of patients. Physical quality of life was important to 70% of patients, and mental quality of life was important to 52% of patients. The rest of the 11 concerns were noted as important by fewer than half of patients (Fig. 1); notably, despite the costs noted above, 79% of patients reported that financial issues were not a concern.

DISCUSSION

Patients with early stage HPV-related oropharyngeal cancer offer insights into ways to optimize surveillance and survivorship care. As the treatment paradigm for HPV-related cancers continues to evolve, efforts must also focus on individualizing follow-up care based upon the outcomes of the disease itself and patient preferences.

Substantial work in careful treatment deintensification in these excellent-prognosis patients is ongoing. Although RTOG 1016 [6] and De-ESCALaTE [38] have shown that de-escalating systemic therapy with cetuximab failed, other

Table 2. Health-related preferences

	<i>n</i> (%)
Self-assessment of physical health	
Excellent	14 (20.9)
Very good	29 (43.3)
Good	18 (26.9)
Fair	5 (7.5)
Trust in health care provider	
Yes, definitely	66 (98.5)
Yes, to some extent	1 (1.5)
No, not at all	0 (0.0)
Shared decision-making preferences	
I prefer to make the final treatment decision	0 (0.0)
I prefer to make the final treatment decision after seriously considering my doctor's opinion	24 (35.8)
I prefer that my doctor and I share responsibility for deciding which treatment is best	33 (49.3)
I prefer that my doctor makes the final treatment decision, but seriously considers my opinion	8 (11.9)
I prefer to leave all treatment decisions to my doctor	2 (3.0)
Medical minimizer-maximizer scale	
Maximizer	48 (71.6)
Minimizer	19 (28.4)
Health literacy: confidence in filling out forms	
All of the time	58 (86.6)
Most of the time	6 (9.0)
Some of the time	1 (1.5)
A little of the time	2 (3.0)
None of the time	0 (0.0)
Worry of cancer recurrence	
Mean	5.48
SD	3.56

approaches are investigating numerous promising approaches to de-escalation such as the use of induction chemotherapy [39], decreasing radiation dose [40] or volumes [41], or modifying indications for postoperative treatment. As these investigations cautiously continue, standard of care therapy should remain unchanged, as standard of care is associated with excellent outcomes. In contrast to treatment studies, current surveillance paradigms do not differentiate follow-up recommendations by HPV status and do not adapt to the excellent outcomes in HPV-related populations. The current study aimed to obtain the patient perspective on altered surveillance to inform strategies to optimize surveillance in the HPV-related population.

A substantial portion of patients (61.2%) would be interested in a post-treatment surveillance option that included fewer clinic visits. There was no significant increase in interest with additional information provided in the vignette for scenario 2, possibly reflecting the high knowledge of this

Table 3. Costs associated with surveillance visits

Dollar amount	Copay, <i>n</i> (%)	Out-of-pocket costs, <i>n</i> (%)	Total costs, <i>n</i> (%)
\$101+	3 (4.5)	6 (9.0)	15 (22.4)
\$51–100	2 (3.0)	9 (13.4)	7 (10.4)
\$26–50	12 (17.9)	13 (19.4)	13 (19.4)
\$1–25	6 (9.0)	27 (40.3)	27 (40.3)
\$0	13 (19.4)	5 (7.5)	5 (7.5)
N/A	31 (46.3)	7 (10.4)	7 (10.4)

Copay costs indicate money due by the patient at the time of appointment. Out-of-pocket costs reflect gas, food, lodging, or other nonmedical costs. Total costs are the sum of these two for all patients that reported at least one of these costs; not available in this latter column indicates participants who did not mark any costs down for either prior column. Of note, costs assessed here did not include medical bills or additional subsequent costs from the health care system or insurers.

population regarding HPV-related cancer outcomes and etiology. Cancer-related knowledge was higher in this study as compared with previous reports, which have previously suggested that the proportion of patients with HPV-related oropharyngeal cancer understanding the viral etiology of their cancer may be as low as 35% [28, 42]. The exploratory analysis of factors that correlated with interest in altered surveillance is in line with what many physicians intuitively know to be true: some patients are intrinsically more likely to seek out health care (medical maximizers) or are more worried about cancer recurrence [43]. In this article, we see that these intuitive perceptions about patients may play out as interest in novel surveillance or treatment paradigms.

It is possible that the proportion of patients who are interested in remote approaches or decreased surveillance may have increased because of the ongoing COVID-19 pandemic and current interest in fewer surveillance visits may be higher under current circumstances. Additionally, telemedicine capabilities have increased as a result of COVID-19 in tandem with favorable reimbursement modifications, likely making remote monitoring more accessible at head and neck cancer centers [44]. A recent telephone-based quality of life survey in patients with oral cavity cancer showed that remote monitoring via patient-reported outcomes may offer an excellent means to detect cancer recurrences while managing appointment-related burdens [45].

In this study, patients reported significant burdens associated with surveillance visits at a tertiary care facility, including a substantial distance driven for each appointment, allotting a significant portion of a working day for a single visit, and costs including copays and other non-medical costs associated with each visit. Interestingly, a substantial portion of patients in this sample reported uncertainty with identifying their typical copay costs (46.3%) and other nonmedical out-of-pocket costs (10.4%), with the former possibly reflecting poor transparency in medical billing and suggesting that these financial burdens may be higher than reported in this study. Despite this, 22.4% of patients reported spending >\$100 per visit on these costs.

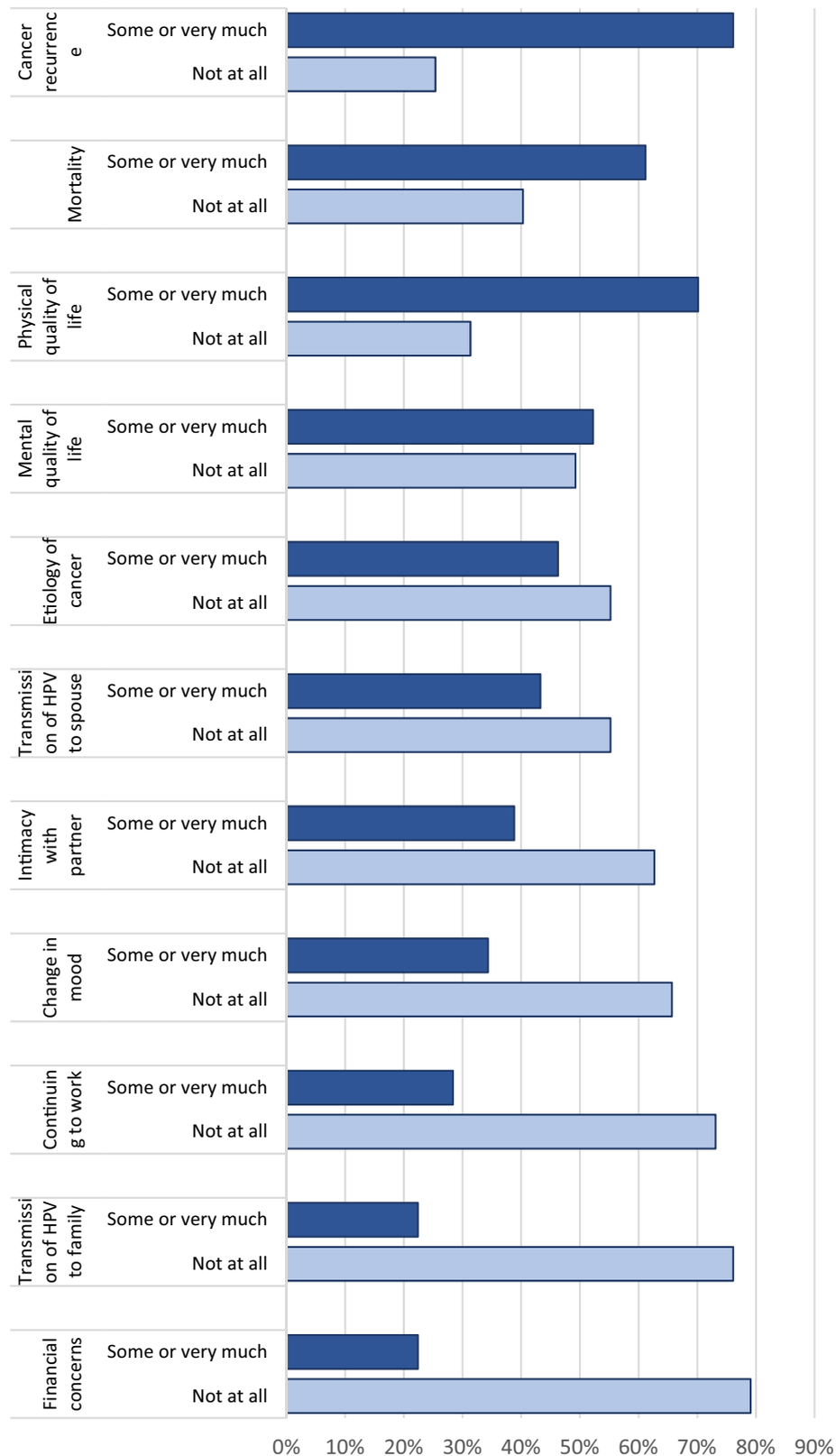


Figure 1. Treatment-related concerns listed as important by patients.

Importantly, almost all patients in this study were insured, and the costs reported in this article likely underestimate these burdens in the wider population. Additionally, these costs did not include direct medical bills related to

surveillance. We have previously shown that 33% of patients with head and neck cancer going through radiation treatment reported at least a moderate financial burden from treatment, and this was associated with increased

treatment noncompliance [46]. Although 79% of participants in this study reported that financial issues were not a concern, possibly reflecting the relatively affluent population studied here, options that incorporate fewer in-person surveillance visits and more remote monitoring may help offset these patient-borne burdens without compromising ability to detect recurrences.

Patients reported concerns that they wished to address during surveillance visits. Consistent with prior reports [37], patients rated cancer recurrence as the most important concern (76%); the next most important concern was physical quality of life (70%). Current surveillance approaches focus on the risk of local recurrence, with frequent nasopharyngoscopy to directly assess for recurrence. Despite clinical practice guidelines that have endorsed comprehensive survivorship care [47, 48], it remains challenging to implement surveillance approaches that cover all of these comprehensive facets and new approaches are needed. For example, incorporation of quality of life and patient-reported outcomes into optimized surveillance approaches may offer a means to meet goals outlined for survivorship care.

Because of recent publications investigating the use of circulating tumor DNA in surveillance of patients with HPV-related cancers [49], this topic was included in the survey. Patients were most interested in including blood tests as a nonclinic-based surveillance option to incorporate into care. This may be reflective of standard association of blood tests with clinical care; it may also be reflective of publicity surrounding the use of circulating tumor DNA to follow patients with HPV-related cancers [49]. Further studies validating the role of circulating tumor DNA into surveillance are needed. When allowed to select multiple options for adjuncts to surveillance, more than half of patients selected urine tests or surveys, offering two additional tools to add to surveillance that could potentially be administered remotely and subsequently minimize surveillance-related burdens. Additionally, patient-reported quality-of-life metrics may offer a means to tailor survivorship symptom-directed visits and better incorporate patient preferences. They may also allow for detection of recurrence, with changes in quality-of-life score potentially predicting for both local and distant recurrence [45, 50]. Systematic administration of quality of life instruments in the metastatic setting has been shown to improve quality of life [51] and increase overall survival [52]. Patients in this study were least interested in symptom-directed visits such as speech-language pathology visits for dysphagia, suggesting that engaging patients in identifying optimal methods to incorporate survivorship care is needed.

Current surveillance recommendations do little to address long-term quality of life or survivorship issues in patients with head and neck cancer. Survivorship care after cancer treatment has several components: the detection of recurrences and new cancers, which is well addressed by current surveillance recommendations, but also physical effects of treatment, psychosocial effects of treatment, health promotion, and management of chronic conditions [53], the latter four of which are poorly addressed by

current surveillance paradigms despite publication of survivorship guidelines [47, 48]. With patient interest in decreased clinic visits as demonstrated in the current survey study and need to address these additional domains of survivorship care, there is opportunity to improve surveillance paradigms especially in this good-prognosis group of patients with HPV-related oropharynx cancer.

Strengths of this study include its high survey response rate, indicating that these responses likely represent the wider early stage HPV population seen at this tertiary academic center. The older white male population is reflective of the population predicted to hold the largest burden of HPV-related cancers in the U.S. in the coming decades [54]. Limitations of this study include its small sample size, single institution, and lack of diversity. Minorities have worse outcomes in this cancer subtype [55], and it may be expected that the characteristics and perspectives described in this study may differ for a more diverse population, such as one that is more diverse in race, income, and insurance status. Additionally, all patients surveyed here were patients at a tertiary care institution, and it is unclear how priorities and burdens may change for patients treated in smaller community centers. Finally, this study did not include items specific to telemedicine or radiologic surveillance for distant disease. Current head and neck cancer guidelines do not incorporate routine imaging, but recent publications suggest this may be of interest in developing future altered surveillance methods [56, 57].

CONCLUSION

Patients with early stage HPV-related oropharyngeal cancers are interested in altered survivorship paradigms, experience significant time and cost burdens related to surveillance visits, and have concerns that are not well addressed in the current paradigm including physical and mental quality of life. Optimized surveillance approaches should incorporate these patient priorities and minimize associated burdens.

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DISCLOSURES

The authors indicated no financial relationships.

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