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Investigating the impact of self-management behaviours on quality of life and fear of recurrence in head and neck cancer survivors: a population-based survey.

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Keywords: cancer, cancer survivorship, fear of recurrence, head and neck cancer, oncology, psycho-oncology, quality of life, self-management.

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ABSTRACT

Objectives: Emerging cancer survivorship research suggests that self-management can lead to improved outcomes. However, research examining the impact of self-management behaviours on quality of life (QoL) and fear of recurrence (FoR) in cancer survivors is lacking. This study investigated the relationship between self-management behaviours and QoL and FoR following treatment for head and neck cancer (HNC).

Methods: Postal surveys were sent to 734 eligible HNC survivors (ICD10 C01-C14;C32) in the Republic of Ireland who were 12-60 months post-diagnosis. QoL and FoR were measured using the Functional Assessment of Cancer Therapy (FACT-G & Head & Neck Cancer Subscale) measure and Fear of Relapse/Recurrence Scale, respectively. Seven self-management behaviours were measured using the Health Education Impact Questionnaire.

Results: Three-hundred-and-ninety-five HNC survivors completed surveys (50.3% response rate). After controlling for socio-demographic and clinical characteristics, self-management behaviours accounted for 20-39.4% of the variance in QoL and FoR. Higher scores on *positive and active engagement in life, constructive attitudes and approaches* and *skill and technique acquisition* were significantly associated with higher global QoL and lower FoR, while higher scores on *positive and active engagement in life* and *constructive attitudes and approaches* only were significantly associated with higher HNC-specific QoL. Additionally, lower scores on *self-monitoring and insight* were significantly associated with higher HNC-specific and global QoL and lower FoR.

Conclusions: The findings highlight the potential utility of self-management interventions promoting active problem-solving, positive self-talk, and skill acquisition among cancer survivors. However, increased self-monitoring may relate to negative outcomes in HNC; a finding which warrants further investigation.

INTRODUCTION

Medical advances in recent decades have increased the number of people who are living longer with cancer and its consequences [1]. Many cancer survivors' daily lives are affected by ongoing physical, emotional and social needs arising from their condition and its treatment [2]. Arising from the enduring nature of these needs, survivors must respond to new and evolving demands and assume an active role in maintaining health and well-being following primary treatment [3]. In this context, cancer self-management has been defined as "awareness and active participation by the person in their recovery, recuperation and rehabilitation, to minimise the consequences of treatment, and promote survival, health and well-being" [4, p.6]. Key self-management behaviours in cancer survivorship include health-directed behaviours like eating a healthy diet [5], monitoring the consequences of cancer and its treatment [3], effectively navigating health services resources [6], adopting constructive attitudes to manage emotional challenges [7] and drawing upon social support [8]. Emerging research suggests that interventions which promote such behaviours in cancer survivorship can "improve symptoms of fatigue, pain, depression, anxiety, emotional distress and quality of life (QoL)" [9, p.1323].

Head and neck cancer (HNC) survivors often have wide-ranging needs requiring ongoing self-management. These include eating and speech difficulties, facial disfigurement and elevated distress, which have marked negative effects on survivors' QoL following treatment [10-12]. Fear of recurrence (FoR) can also be a persistent and significant issue in this population, with noted deleterious effects on QoL [12-13]; FoR is the most frequently cited concern by HNC survivors aged less than 70 years [14]. Our recent qualitative research has illustrated the variety of strategies that HNC survivors use to self-manage these ongoing issues [15], the barriers they encounter in their self-management attempts [16], and the processes through which they integrate self-management into their daily lives [17]. However, to our knowledge, there has been no research on the relationship between self-management behaviours and QoL in this population to date, nor has there been any research on the relationship between self-management behaviours and FoR in cancer more generally.

Understanding the relationship between self-management and these key outcomes in cancer

survivorship is important in order to inform the content and delivery of self-management interventions designed to develop the skills and confidence that cancer survivors need to live well following treatment. The aim of the current paper was to examine the degree to which HNC survivors' performance of self-management behaviours were associated with their QoL and FoR in the post-treatment period, after controlling for key demographic and clinical characteristics. We hypothesized that self-management behaviours would be associated with higher QoL and lower FoR among HNC survivors.

MATERIALS AND METHODS

Design, participants and recruitment procedures

We adopted a national population-based postal survey design targeting HNC survivors in the Republic of Ireland. This population-based survey is part of a larger study to develop a self-management intervention to improve quality of life among HNC survivors. We received ethical approval from 20 hospitals across the country¹, including all major treatment centres for HNC. Between May-June 2015, we identified 1,208 individuals eligible for the study in collaboration with the National Cancer Registry Ireland (NCRI), which records all new primary cancers diagnosed among people residing in the country. The study population was identified from the NCRI using the following criteria to indicate that an individual: 1) was alive, 2) was aged 18 years or older at diagnosis, 3) had obtained a diagnosis of a primary invasive tumour in the head and neck region, including cancers of the oral cavity, oropharynx, larynx, hypopharynx, paranasal sinuses, and salivary glands (ICD10 C01-C14, C32), 4) was 1-5 years post-diagnosis (to ensure post-treatment status), 5) had an absence of a second invasive cancer diagnosed after HNC (excepting non-melanoma skin cancer), and 6) had been treated in one of the 20 participating hospitals. We then identified 82 consultants who had treated these

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survivors and requested their assistance to screen them to confirm eligibility. Twenty-seven consultants, accounting for 125 HNC survivors, either did not agree or failed to respond to our screening request. The remaining 55 treating consultants (with 1,083 eligible patients) agreed to screen a list of their patients and confirm eligibility. These consultants confirmed that an individual 1) was alive, 2) was aware they had cancer, 3) was not receiving palliative care, 4) had not developed a second invasive cancer (apart from non-melanoma skin cancer), 5) had completed primary treatment for HNC, 6) was not receiving treatment for a recurrence or secondary cancer, 7) was considered disease-free for the preceding four months, and 8) there was no medical or other reason why it would be inappropriate to contact them about the study. Following this screening process, survey packs including a survey booklet, explanatory letter, information sheet, consent form and stamped-addressed return envelope, were posted to 785 eligible individuals between November 2015-March 2016. Non-responders received up to two reminder letters (at 2-4 weeks after issue of the survey pack).

Supplementary Figure 1 further illustrates the screening/recruitment process for interested readers.

Participants were asked to indicate if they received help filling in the questionnaires (e.g. because of poor sight, difficulty writing, etc.). The survey, which we anticipated would take approximately 30 minutes to complete, contained ten sections; only data pertaining to demographics, self-management behaviours, QoL and FoR are reported in the present paper.

Measures

We used the Health Education Impact Questionnaire [HEIQ] to determine participants' use of self-management behaviours. There is currently an absence of high-quality instruments to measure self-management behaviours. However, this measure has been extensively used for this purpose across a broad range of chronic conditions and was devised by Australian researchers to assess the efficacy of patient education programs in teaching key self-management behaviours [18]. The measure asks respondents to indicate their level of agreement with a series of statements across eight domains (42-items) on four-point Likert scales and has been used as a measure of self-management behaviours with other groups of cancer patients [19-20]. Higher scores in seven domains indicate greater

performance of specific self-management behaviours (*positive and active engagement in life, health-directed behaviour, constructive attitudes and approaches, skill and technique acquisition, self-monitoring and insight, health service navigation, and social integration and support*). Higher scores on the *emotional well-being* domain denote higher levels of negative affect.

We assessed QoL using the Functional Assessment of Cancer Therapy (FACT-G). This consists of 27 statements, which ask participants to consider the previous week and subsequently rate their QoL across physical, social/family, emotional and functional domains on a five-point scale. We also used the FACT HNC-specific domain; the Head and Neck Cancer Scale (HNCS). This comprises 12 statements about HNC-specific challenges, and can be used independently of the FACT-G to indicate HNC-specific QoL. FACT-G and HNCS scores were computed for all individuals based on the authors' scoring instructions. In all cases, higher domain scores indicate higher QoL. The FACT-G and HNCS have been widely used and validated in different countries [21-22] and are commonly used to assess the QoL of HNC survivors [12].

We assessed FoR using the Fear of Relapse/Recurrence Scale [FRRS; 23]. On this measure, respondents are asked to rate five statements on a five-point Likert scale as they apply to the previous week. Scores range from 5-25 with higher scores indicative of higher levels of FoR. Thewes et al. [24] have found that this scale is the most extensively validated brief measure of FoR with desirable psychometric properties.

Sociodemographic and clinical data was collected via NCRI data and survey data. We obtained NCRI data on participants' age (in years) at time of survey completion, sex, cancer site (four variables categorized as larynx, salivary glands, oral cavity or pharynx, respectively), treatment received within the first year since diagnosis (three variables categorized as whether not the individual had received cancer-directed surgery, radiotherapy or chemotherapy) and cancer stage at diagnosis. Survey data was used to indicate participants' smoking status (current smoker or non-smoker), relationship status (in a relationship or not), education level (primary, secondary, third level/postgraduate or unknown),

presence of a comorbidity (at least one or none), and alcohol use (extent of use, assessed using the Alcohol Use Disorders Identification Test [AUDIT-C; 25], which measures the extent of problem drinking behaviours across three items; scores above four indicate that an individual's drinking may be negatively affecting their safety).

Statistical Methods

In the following analyses, we only included the 7 domains of the HEIQ relating to self-management behaviours (i.e. we excluded *emotional well-being* as it is a measure of negative affect). Cronbach's alpha values for all included scales and associated subscales were between 0.83-0.94. For respondent/non-respondent analysis, we used logistic regression with key sociodemographic and clinical characteristics aggregated from survey and NCRI data as predictor variables. We used hierarchical multiple regression analysis to explore associations between self-management behaviours and global QoL, HNC-specific QoL and FoR scores, respectively, after controlling for relevant sociodemographic and clinical characteristics.

In preparation for hierarchical multiple regression analyses, we explored the relationships between key sociodemographic and clinical characteristics (age at diagnosis, sex, self-reported current smoking status, extent of alcohol use, presence of a comorbidity, treatment received, cancer location, cancer stage at diagnosis, employment status, relationship status, number of months between diagnosis and survey completion and whether or not a respondent was helped to complete the survey) and outcome variables through relevant tests of associations (e.g. correlations, t-tests and chi-squares). This exploratory analysis revealed that age at diagnosis, current smoking status, extent of alcohol use and comorbidity presence had significant relationships with global QoL and FoR, and were retained as covariates in multivariate analyses. These same four variables, in addition to three binary predictors relating to cancer type (larynx, salivary glands and pharynx), also had significant relationships with HNC-specific QoL; all seven were retained as covariates for multivariate analysis. In each model, we assessed the independent effects of self-management behaviours on our outcomes by entering pertinent sociodemographic and clinical covariates in the first block, and self-management behaviours

in the second block. We made the a priori decision to exclude the HEIQ subscale *social integration and support* from global QoL multivariate analyses due to potential multi-collinearity with the Social Well-Being subscale, which contributes to FACT-G scores. Although there are subtle differences between these two subscales in how they deal with social interactions (the former specifically relates to receiving help for one's health-related issues, while the latter relates to one's sense of social well-being more generally), we were concerned that the inclusion of the *social integration and support* subscale would inflate the error variance in our analysis of global QoL.

All tests were two-tailed with an alpha level set at 0.05. All statistical analyses were performed using IBM Statistics Package for the Social Sciences, Version 23.

RESULTS

Sociodemographic and clinical characteristics for participants are provided in Table 1. Of the 785 individuals invited to participate, 395 completed and returned surveys (50.32% response rate). Ninety-five participants (24.1%) required help in completing the survey. Logistic regression analysis revealed no significant differences between respondents and non-respondents on the following characteristics (aggregated from survey and NCRI data): age at diagnosis, sex, current smoking status, treatment received, cancer type, cancer stage or time since diagnosis.

[Insert Table 1]

Prior to multivariate analyses, preliminary analyses confirmed that the data did not violate any assumptions of multiple regression. Relevant tests of association for all variables included in the analyses are provided in Supplementary Table 1. All seven self-management behaviours were correlated with each other to a highly significant degree ($p < .001$ in each case).

When we entered sociodemographic and clinical covariates (age at diagnosis, self-reported current smoking status, extent of alcohol use and presence of a comorbidity) in the first block of our global QoL model ($M=80.81$, $SD=18.58$), they accounted for 11.7% of the variance [$F(4,306)=10.17$, $p < 0.001$]. The six self-management behaviours entered in block 2 accounted for an additional 39.4%

of variance in QoL; R^2 -change=0.394, F -change(6,300)=40.27, $p<0.001$. The final model accounted for 51.1% of variance in QoL scores; $F(10,300)=31.36$, $p<0.001$. The following self-management variables independently predicted global QoL: *positive and active engagement in life, self-monitoring and insight, constructive attitudes and approaches* and *skill and technique acquisition*. Of these, *positive and active engagement in life* ($B=17.27$, $CI=12.67-21.87$, $p<.001$) and *constructive attitudes and approaches* ($B=9.84$, $CI=5.85-13.82$, $p<.001$) had the strongest positive association with global QoL, while *self-monitoring and insight* was the only self-management behaviour to have a negative association with global QoL ($B=-9.23$, $CI=-15.13--3.33$, $p<.01$).

[Insert Table 2]

For our HNC-specific QoL ($M=27.76$, $SD=8.46$) model, the socio-demographic and clinical covariates accounted for 15.1% of variance in HNC-specific QoL scores [$F(7,303)=7.71$, $p<0.001$].

The seven self-management behaviours entered in the second block of this model explained an additional 20% of variance; R^2 -change=0.20, F -change(7,296)=13.01, $p<0.001$. The final model accounted for 35.1% of the variance in HNC-specific QoL; $F(14,296)=11.43$, $p<0.001$. The following self-management variables independently predicted HNC-specific QoL: *positive and active engagement in life, self-monitoring and insight* and *constructive attitudes and approaches*. *Positive and active engagement in life* had the strongest positive impact on HNC-specific QoL ($B=5.73$, $CI=3.26-8.20$, $p<.001$); *self-monitoring and insight* was the only self-management behaviour to have a negative relationship with HNC-specific QoL ($B=-5.17$, $CI=-8.32--2.01$, $p<.01$).

In relation to FoR ($M=13.74$, $SD=4.57$), block 1 variables accounted for 7.2% of variance [$F(4,306)=5.94$, $p<0.001$]. The seven self-management behaviours accounted for an additional 32.9% of variance; R^2 -change=0.329, F -change (7,299)=23.49, $p<0.001$. The final model accounted for 40.1% of variance in FoR; $F(11,299)=18.22$, $p<0.001$. The following self-management variables independently predicted FoR: *positive and active engagement in life, self-monitoring and insight, constructive attitudes and approaches* and *skill and technique acquisition*. *Positive and active engagement in life* ($B=-3.18$, $CI=-4.45--1.90$, $p<.001$) and *constructive attitudes and approaches* ($B=-$

2.34, CI=-3.52--1.17, $p<.001$) were most strongly associated with lower FoR, while *self-monitoring and insight* was the only self-management behaviour to be associated with increased FoR ($B=2.01$, CI=.39-3.63, $p<.05$) among participants. Further details on the models are provided in Table 2.

CONCLUSIONS

To our knowledge, this is the first study to investigate the relationship between key self-management behaviours and QoL and FoR in HNC survivors, and the first to investigate the relationship between self-management behaviours and FoR in cancer more generally. Our results demonstrate that higher scores on three key self-management behaviours (*positive and active engagement in life, constructive attitudes and approaches* and *skill and technique acquisition*) were significantly associated with higher global QoL and lower FoR among HNC survivors who had completed their active treatment. In contrast, higher *self-monitoring and insight* were significantly associated with lower levels of QoL and higher FoR in our sample. The global QoL findings were also replicated with HNC-specific QoL, with the exception of a non-significant association between *skill and technique acquisition* and HNC-specific QoL.

Our findings that *positive and active engagement in life, constructive attitudes and approaches* and *skill and technique acquisition* were significantly associated with increases in global QoL and decreases in FoR, and that *positive and active engagement in life* and *constructive attitudes and approaches* were associated with increases in HNC-specific QoL, are consistent with the wider cancer self-management literature. For instance, positive appraisals such as optimism, hope and inner strength have been linked to positive outcomes such as benefit-finding, post-traumatic growth and higher QoL in individuals with HNC [7; 26], and greater psychosocial adjustment and lower perceived stress among cancer survivors more generally [27-28]. Engaging in active problem-solving is a core component of cancer self-management interventions [29], while acquiring new skills and techniques (e.g. relating to medical management of HNC site) is recognised as critical in HNC rehabilitation [30]. Importantly, our findings indicate that these self-management behaviours are strongly associated with improved QoL and reduced FoR among HNC survivors, even after

controlling for pertinent socio-demographic and clinical factors. In line with these findings, HNC survivors might benefit from structured self-management interventions with components which focus on active problem-solving, positive appraisal and acquiring specific skills and techniques that are relevant to self-managing their condition (while a thorough discussion of such interventions is beyond the scope of the current paper, interested readers are directed to Davies & Batehup's [4] review on this topic).

An additional striking finding in this study was that increased levels of *self-monitoring and insight* were associated with poorer QoL and FoR outcomes in HNC survivors. While these findings were contrary to our hypotheses relating to self-monitoring, they are perhaps unsurprising, given that elevated levels of self-monitoring may be closely linked to hypervigilance to signs of recurrence in cancer survivors. Hypervigilance is a key component in conceptualisations of FoR [31] and maladaptive coping in HNC [32-33], and is associated with the incidence of post-traumatic stress disorder in cancer survivorship [34-35]. In contrast, self-monitoring has typically been regarded as adaptive for individuals with cancer [3], with recommendations to target this behaviour in cancer self-management interventions [9] and qualitative accounts supporting its positive effects [36]. In light of this, further research is needed to gain a better understanding of the impact of self-monitoring on outcomes in HNC survivorship; e.g. to establish if a threshold point exists where adaptive self-monitoring becomes maladaptive hypervigilance in HNC survivors. Furthermore, self-management behaviour scales like the HEIQ may need to be revised for use in cancer survivors generally in order to account for the potential continuum of adaptive self-monitoring to maladaptive hypervigilance in cancer survivorship.

We also found that higher scores on *health-directed behaviours, health service navigation and social integration and support* were associated with higher QoL and lower FoR among HNC survivors in univariate analyses, but these associations did not remain significant after controlling for relevant socio-demographic and clinical characteristics. This suggests that the variance in QoL and FoR attributable to these self-management behaviours may have been accounted for by our covariates, or

may have been confounded by the presence of other self-management behaviours with which they were highly correlated. In spite of this, extant research has demonstrated the importance of these behaviours in cancer self-management [5-6; 8]. With this in mind, further research is needed to determine the extent to which these behaviours benefit cancer survivors across various stages of the cancer trajectory and/or in relation to other important outcomes (e.g. 5-year survival). Furthermore, due to their potential shared variance with other variables, more refined measures of these self-management behaviours may be required which better capture the potential unique variance that they account for in relation to important outcomes for HNC survivors (and, potentially, in relation to other cancer types). Indeed, Schuler et al.'s [37] work has indicated that this instrument may need to be adapted for specific conditions like HNC.

Study Limitations

The current study was cross-sectional; as such, we cannot be certain of the direction of associations between self-management behaviours, QoL and FoR. Future longitudinal research is needed to establish this, and whether or not there may be different relationships between HNC survivors' use of self-management behaviours and their QoL and FoR at different stages in the post-treatment trajectory. Our sample also comprised HNC survivors with a wide range of time since the end of their treatment. Emerging research suggests there are core differences between HPV and non-HPV-related HNC survivors in their QoL [38]; NCRI data relating to HPV was not available to us for HNC cancer survivors at the time of data collection, so we could not explore this. Additionally, 27 consultants (accounting for approximately 10% of HNC survivors meeting our eligibility criteria in Ireland) did not take part in our study. We were also unable to adjust for consultants or treating hospitals in our analyses as the NCRI data available to us was limited to the physician and hospital site registered at diagnosis (and these may not be the centre/clinician responsible for the survivor's cancer care); these variables may have accounted for some of our unexplained variance. Nonetheless, the rigorous population-based survey design was a clear strength, ensuring the sample was broadly reflective of HNC survivors in the Irish population following completion of their primary treatment when compared with data from the NCRI database. Furthermore, to our knowledge, this is the first study to

investigate the relationship between key self-management behaviours and QoL and FoR in HNC survivors, and the first to investigate the relationship between self-management behaviours and FoR in cancer more generally; our findings clearly demonstrate the positive association between key self-management behaviours and these important outcomes.

Clinical Implications

The current findings suggest that developing effective ways of supporting self-management behaviours like active problem-solving and positive self-talk [27; 39] may lead to increases in QoL and decreases in FoR among HNC survivors following treatment. The findings also have important implications for the design and implementation of supported self-management interventions for HNC survivors. Adjustment-focused self-management interventions supporting the acquisition of specific self-management skills [36] may lead to positive effects in relation to QoL and FoR for HNC survivors. Furthermore, self-management intervention components related to self-monitoring may need to be specifically tailored to consider the potential for maladaptive self-monitoring among HNC survivors (and, potentially, among other cancer survivors). Health professionals should also be cognisant of the potential negative impact of this behaviour when advising HNC survivors on recognising and managing the signs and symptoms of recurrence and the development of specific screening tools to capture maladaptive hypervigilance may be useful for healthcare professionals to use in the clinical setting. Similarly, routine screening for maladaptive FoR (e.g. using the Patient Concerns Inventory [40]) may help to identify those who need to be targeted with structured supported self-management programmes.

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CONFLICT OF INTEREST

There is no conflict of interest to report.

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Table 1: Socio-demographic/clinical characteristics of participants

<i>Variable</i>	<i>N</i>	<i>Variable</i>	<i>N</i>	<i>Variable</i>	<i>N</i>
Total	395				
Sex		Time since diagnosis		Self-reported smoking status	
Male	272	12-24 months	125	Current smoker	69
Female	123	25-36 months	98	Ex-smoker	221
Relationship status		37-48 months	95	Never smoked	90
Married/cohabiting	248	49-60 months	77	Unknown	15
Not in a relationship	132	Cancer location		Age	
Unknown	15	Larynx	119	25-39 years	18
Treatment(s)		Salivary glands	24	40-54 years	104
Chemotherapy only	4	Oral cavity	143	55-69 years	197
Radiotherapy only	64	Pharynx	110	70 years or more	76
Surgery only	109	Cancer stage at diagnosis		Highest level of education	
Radiotherapy/chemotherapy	54	I	120	Primary	114
Surgery/chemotherapy	2	II	38	Secondary	187
Surgery/radiotherapy	64	III	44	Third level/ Postgraduate	89
Surgery/radiotherapy/chemotherapy	30	IV	103	Unknown	5
Unknown	68	Unknown	90		
Employment status		Presence of a comorbidity			
Currently working	110	Yes	196		
Not working/Retired	280	No	196		
Unknown	5	Unknown	3		

Table 2. Summary of hierarchical multiple regression models predicting global QoL, HNC-specific QoL and FoR

Global QoL						HNC-specific QoL								
Variable		95% CI				Variable	95% CI				Variable			
	B	SE	β	Lower	Upper	R ²	B	SE	β	Lower	Upper	R ²		
Step 1						0.12***	Step 1					0.15***	Step 1	
Step 2						0.51***	Step 2					0.35***	Step 2	
(Constant)	5.56	7.59		-9.38	20.51		(Constant)	6.33	4.08		-1.69	14.35		(Constant)
Age	.27***	.07	.16	.13	.40		Age	.08*	.04	1.00	.00	.15		Age
CS	-3.93	2.02	-.08	-7.91	.06		CS	-.91	1.07	-.04	-3.02	1.20		CS
Alc	.46	.31	.06	-.16	1.07		Alc	.55**	.17	.16	.22	.88		Alc
Comorb	-3.97*	1.56	-.11	-7.05	-.91		Comorb	-1.18	.83	-.07	-2.81	.44		Comorb
HDB†	-.28	1.46	-.01	-3.16	2.59	R ²	Larynx	1.89	1.00	.10	-.09	3.86		HDB†
						change:								
PAE†	17.27**	2.34	.48	12.67	21.87	.39***	Salivary glands	6.76***	1.76	.19	3.30	10.21		PAE†
SMI†	-9.23**	3.00	-.19	-15.13	-3.33		Pharynx	-1.88	1.01	-.10	-3.87	.11		SMI†
CAA†	9.84***	2.03	.29	5.85	13.82		HDB†	-.29	.77	-.02	-1.81	1.24	R ²	CAA†
													change:	
STA†	5.26*	2.46	.14	.42	10.09		PAE†	5.73***	1.26	.35	3.26	8.20	0.20***	STA†
HSN†	-2.72	2.26	-.07	-7.18	1.73		SMI†	-5.17**	1.60	-.23	-8.32	-2.01		SIS†
							CAA†	3.80**	1.16	.25	1.51	6.09		HSN†
							STA†	1.85	1.31	.10	-.74	4.44		
							SIS†	-.24	1.17	-.02	-2.54	2.06		
							HSN†	-.49	1.26	-.03	-2.97	1.99		

Key: *($p < .05$); **($p < .01$); ***($p < .001$); †HEIQ subscale

HDB=Health-Directed Behaviour; PAE=Positive and Active Engagement in Life; SMI=Self-Monitoring and Insight;

CAA=Constructive Attitudes and Approaches; STA=Skill and Technique Acquisition; SIS=Social Integration and Support;

HSN=Health Service Navigation; CS=Current Smoker; Alc=Extent of alcohol use; Comorb=Presence of comorbidity