

BMJ Open Women's acceptability of and experience with primary human papillomavirus testing for cervix screening: HPV FOCAL trial cross-sectional online survey results

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ABSTRACT

Objective To study participant's acceptability of and attitudes towards human papillomavirus (HPV) testing compared with cytology for cervical cancer screening and what impact having an HPV positive result may have in future acceptability of screening.

Design Cross-sectional online survey of clinical trial participants.

Setting Primary care, population-based Cervix Screening Program, British Columbia, Canada.

Participants A total of 5532 participants from the HPV FOCAL trial, in which women received HPV and cytology testing at study exit, were included in the analysis. Median age was 54 years. The median time of survey completion was 3 years after trial exit.

Outcome measures Acceptability of HPV testing for primary cervical cancer screening (primary); attitudes and patient perceptions towards HPV testing and receipt of HPV positive screen results (secondary).

Results Most respondents (63%) were accepting of HPV testing, with the majority (69%) accepting screening to begin at age 30 years with HPV testing. Only half of participants (54%) were accepting of an extended screening interval of 4–5 years. In multivariable logistic regression, women who received an HPV positive screen test result during the trial (OR=1.41 95% CI 1.11 to 1.80) or were older (OR=1.01, 95% CI 1.00 to 1.02) were more likely to report HPV testing as acceptable.

Conclusions In this evaluation of acceptability and attitudes regarding HPV testing for cervix screening, most are accepting of HPV testing for screening; however, findings indicate heterogeneity in concerns and experiences surrounding HPV testing and receipt of HPV positive results. These findings provide insights for the development of education, information and communication strategies during implementation of HPV-based cervical cancer screening.

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INTRODUCTION

It is well established that persistent infection with an oncogenic strain of the human papillomavirus (HPV), the most common

STRENGTHS AND LIMITATIONS OF THIS STUDY

- ⇒ Measures of acceptability and patient perceptions of human papillomavirus (HPV) testing within a primary cervix screening programme.
- ⇒ Reports on acceptability of increased screening interval and delayed onset of screening initiation, and impact of a positive HPV test result.
- ⇒ Recommendations for key health promotion messaging to address potential barriers to HPV testing for primary cervical screening.
- ⇒ Limitations include that participants were recruited from a large clinical trial on HPV testing for cervical cancer screening and may not be representative of the general screening population.

sexually transmitted infection (STI) around the world, is the causative agent for most cervical cancers.^{1 2} There is a robust body of evidence regarding the superior performance of HPV versus cytology screening in detection of cervical intraepithelial neoplasia grade two or worse and greater protection against cervical cancer.^{3–5} As such, several countries around the world have implemented primary HPV testing for cervical cancer screening, including Australia, the Netherlands and the UK, with many other jurisdictions in various planning stages for HPV-based screening implementation. A shift to an HPV-based screening approach results in different programme guidelines and, thus, a different experience for the person undergoing screening. The very high negative predictive value of HPV testing permits the interval between screens to be extended to 5 or more years compared with cytology testing, recommended every 2–3 years in most jurisdictions.^{5–7} Due to high prevalence and regression rates of HPV infection in younger women, HPV-based screening may

not be recommended until 25–30 years of age.^{5,6} In addition, being screened for cervical cancer with a test for an STI can result in anxiety and concern for those undergoing screening.^{8,9}

With such a transformative change in what is arguably a well-established screening paradigm, it is crucial to examine women's readiness or acceptance of HPV testing compared with cytology testing for screening to ensure engagement in screening is not hampered by a change in technology or guidelines. This unintended consequence was illustrated in Australia, prior to the change in the national programme from cytology to HPV screening, when a 2017 petition opposing the changes garnered 70 000 signatures.¹⁰ Respondents to the Australian survey indicated concerns about such things as the extended interval and missing cancer cases in younger women as a result of the programme change.¹⁰ Despite enhancements to screening efficacy and safety, a successful change in technology requires acceptance by those who undergo screening. Anticipating women's questions and concerns prior to implementation of programme changes can mitigate resistance to change and assist in the design of targeted education strategies.

This analysis is of the 48-month exit survey for the Human Papillomavirus For Cervical Cancer Screening Trial (HPV FOCAL). HPV FOCAL is currently the only North American trial comparing primary HPV testing to cytology (liquid based) for screening within an organised programme, which also provides us with the unique opportunity to assess women's experiences with HPV testing in a population-based programme setting. The primary objective of this analysis was to explore participant's acceptability of and attitudes towards HPV testing compared with cytology for cervical cancer screening and what impact having an HPV positive result may play in future acceptability of screening.

METHODS

Participants

Survey participants were recruited through the HPV FOCAL trial, a publicly funded randomised control trial comparing primary HPV testing every 4 years (HPV arm) to liquid-based cytology testing (cytology arm) every 2 years for cervical cancer screening. HPV FOCAL recruited women, 25–65 years of age, from two largely metro areas in British Columbia who were due for cervical cancer screening from 2008 to 2012. Participants of the HPV FOCAL trial were engaged in cervical cancer screening through a large population-based screening programme and representative of women at average risk of cervical cancer in North America.¹¹ Trial design and primary outcome results have been previously described in detail.^{3, 11–14} Participants were provided with information on HPV, HPV testing (including differences between Pap and HPV testing and the reasons behind an extended interval between negative HPV screens) and cervical cancer on enrolment and throughout the trial follow-up

period. A total of 9552 women were randomised to the HPV arm and 9457 women to the cytology arm. Women from both the HPV and cytology arms completed trial exit screening between 2012 and 2016, where they received HPV and cytology co-testing at the exit screen. Results were provided to their primary care provider, who then conveyed them to the participants. From August 2017 to February 2018, women from both arms who had attended the 48-month exit screen were invited to complete the online exit survey (figure 1).

HPV FOCAL exit survey

The survey included 26 items that asked participants about HPV knowledge and information seeking before and during the study, acceptability of HPV testing, willingness to increase the screening interval, commencement screening age for HPV testing, attitudes and concerns about test positive results and communication needs around screening results, in addition to demographic details (online supplemental file). Responses included 7-point and 5-point Likert scales, and survey responses were linked to HPV FOCAL trial screen test results. Survey items were based on previous HPV FOCAL surveys assessing HPV testing acceptance.¹⁵ The survey was distributed and managed using the web-based platform of Fluidsurveys (www.fluidsurveys.com). The survey was pilot tested and revised for face and content validity with approximately 20 women, aged 30 years and above prior to distribution to FOCAL participants.

Patient and public involvement

Patient concerns and questions raised during the trial period identified the need for the study, but patients were not involved in the construction of the survey.¹⁶ However, a sample of women who undergo cervical cancer screening in BC were involved in pilot testing of the survey for the purposes of face validity and survey flow and logistics. Based on this feedback, revisions were made to the survey to clarify wording of questions and format of layout.

Response rate and inclusion criteria

Participants from the HPV FOCAL trial from both the HPV and cytology arms who had completed their study 48-month exit screen, had indicated consent to be contacted for future research and for whom email addresses were available were eligible for survey invitation. The invite to complete the survey was sent via email, with one reminder sent a month later for those who had not initiated or completed the survey. Participants were provided with a unique study identifier to access the survey and no personal identifiers were captured during survey completion. Participants were informed they had the option to complete none, some, or all of the survey with completion of survey questions as indication of consent.

Survey completeness was reviewed, and duplicate surveys, where the same woman completed all or some

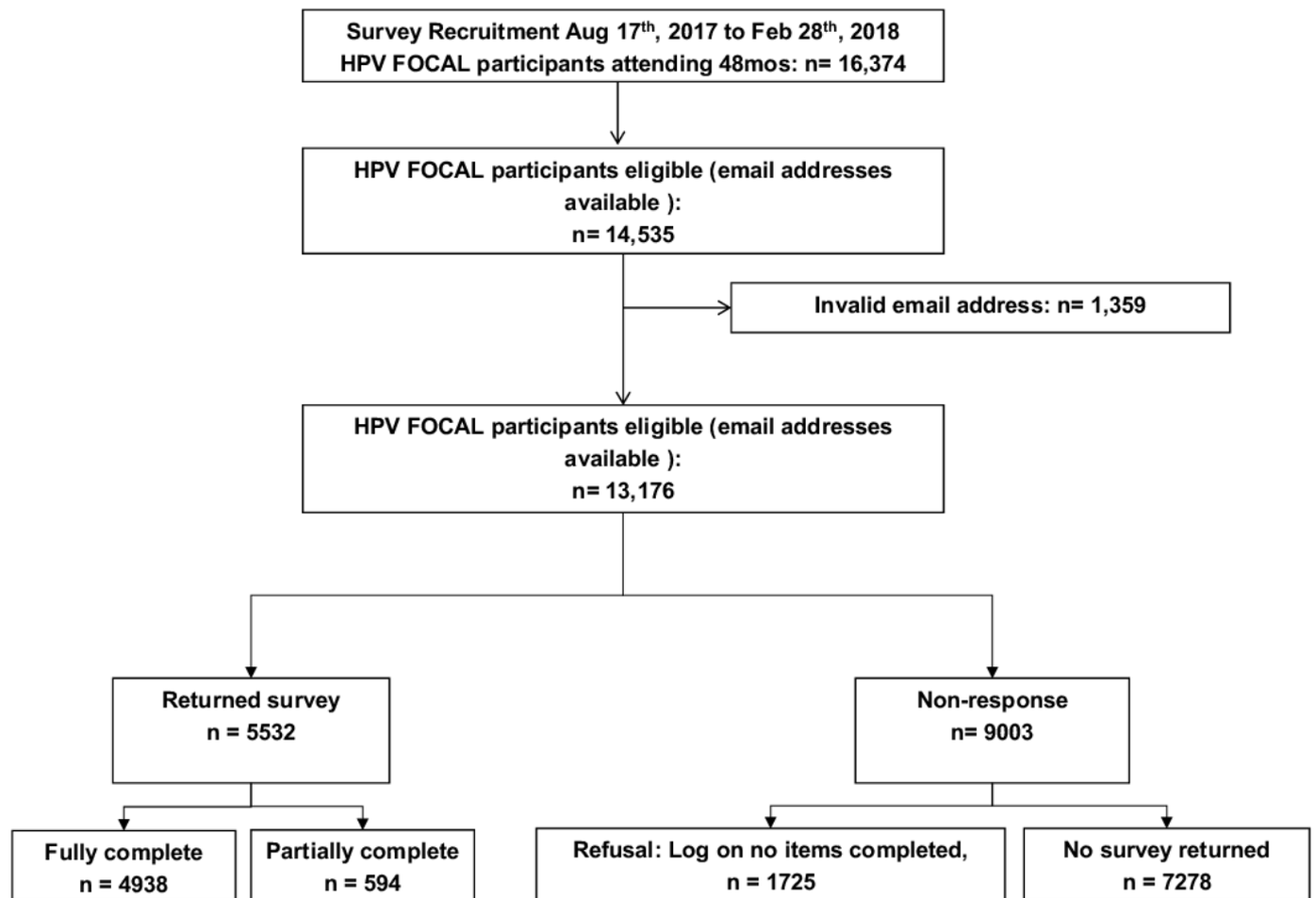


Figure 1 Study flow chart and participant disposition. HPV FOCAL, Human Papillomavirus For Cervical Cancer Screening Trial.

of the survey more than once, were identified. For those with a duplicate entry, the first complete survey was used in the analysis with all other survey attempts discarded. Response rate (%) was the sum of completed surveys plus partial surveys, divided by the number of invitations sent to eligible valid email addresses, as per the American Association for Public Opinion Research guidelines.¹⁷ Non-response included: refusals (clicked the survey link, but did not complete consent or any items), and those assumed eligible with no response received. Email addresses that were undeliverable were considered invalid and not included in the analysis.

Statistical analysis

Survey respondents were compared with HPV FOCAL trial participants on age, study arm, and location of trial recruitment to explore if survey sample was representative of FOCAL trial population.

Our main outcome of HPV testing acceptability was assessed by response to the question ‘having an HPV test instead of a Pap to screen for cervical cancer is acceptable to me’, which was dichotomised from a 5-point Likert scale, with those reporting strongly agree or agree, categorised as ‘accepting’, and those reporting neutral, don’t know, disagree and strongly disagree as ‘not accepting’ of

HPV testing. This categorisation was chosen to capture those who were truly accepting and biased towards the null. Only complete surveys were included, with those who were missing or preferred not to answer excluded. Participants were classified as being HPV positive, if they received an HPV positive screening result at any point during their participation in the HPV FOCAL trial; otherwise, a participant was classified as HPV negative.

Bivariable analysis explored differences in acceptability of HPV testing based on demographic factors such as age, income and education, in addition to HPV screening test result, and length of time since study exit. Factors shown to be potentially associated with acceptability, such as HPV screening starting at 30 years of age and increased screening interval, were also examined.^{16 18 19}

Sociodemographics and attitudes towards HPV testing were explored descriptively with χ^2 for categorical variables and median score test for continuous variables.

Multivariate logistic regression was used to explore the association of the acceptability of HPV testing with a priori identified confounding variables that reached $p \leq 0.2$ in bivariable analysis. Level of significance was 0.05. All statistical analyses were performed in SAS V.9.4 and R V.4.02.

**Table 1** Demographic characteristics of respondents and bivariable analysis of acceptability of HPV testing

		Having an HPV test instead of a Pap smear to screen for cervical cancer is acceptable to me			
		Total (N)	Not acceptable (disagree/neutral/don't know), n (%)	Acceptable (agree), n (%)	P value
Total (N)		5336	1993 (37.4)	3343 (62.6)	
Age	Median (IQR)	5336	53.35 (45.35–61.19)	54.23 (45.64–61.72)	0.051
Marital status†					
	Living with a partner	3806	1438 (78.7)	2368 (76.6)	0.095
	Living without a partner	1115	390 (21.3)	725 (23.4)	
Education†					
	Complete college or higher	3317	1250 (68.5)	2067 (66.7)	0.221
	Incomplete postsecondary or less	1607	576 (31.5)	1031 (33.3)	
Income†					
	Less than \$75 000	1501	529 (32.0)	972 (34.0)	0.175
	\$75 000 or more	3014	1126 (68.0)	1888 (66.0)	
HPV testing status during FOCAL trial					
	Never tested HPV positive	4937	1867 (93.7)	3070 (91.8)	0.015*
	At least one HPV positive result	399	126 (6.3)	273 (8.2)	
Time since exit from FOCAL trial, years					
	Median (IQR)	5336	3.09 (2.26–3.91)	3.04 (2.23,–3.93)	0.616
I would be willing to have an HPV test every 4–5 years instead of a Pap every 3 years†					
	Agree	2858	386 (19.5)	2472 (74.2)	<0.001*
	Disagree	1096	744 (37.6)	352 (10.6)	
	Neutral	1353	847 (42.8)	506 (15.2)	
Receiving HPV testing starting at age 30 years is acceptable to me†					
	Agree	3635	944 (47.8)	2691 (81.0)	<0.001*
	Disagree	682	423 (21.4)	259 (7.8)	
	Neutral	981	608 (30.8)	373 (11.2)	
If cervical cancer screening was to occur every 4 or 5 years, instead of every 3 years, I would be less likely to visit my healthcare provider for other health reasons.					
	Agree	1062	405 (20.4)	657 (19.7)	0.814
	Disagree	3517	1303 (65.7)	2214 (66.4)	
	Neutral	735	274 (13.8)	461 (13.8)	

*Significant to $p < 0.05$.

†Missing values up to 5336=not reported or prefer not to answer.

HPV, human papillomavirus; HPV FOCAL, Human Papillomavirus For Cervical Cancer Screening Trial.

RESULTS

Survey invites were administered from August 2017 to February 2018. A total of 14 535 participants from both the HPV and cytology arms in HPV FOCAL trial were identified as eligible to receive 48-month exit online survey, of which 13 176 were delivered to a valid email address (figure 1). There were 5532 surveys completed, of which 4938 were fully and 594 partially completed.

Characteristics of respondents

The median age of participants completing the survey was 54 years (IQR: 46–62) (table 1). The median time

of survey completion was 3 years after study exit. The majority of respondents (67%) had completed college or higher education and 77% reported living with a partner. Survey respondents were comparable with HPV FOCAL trial participants based on study arm, age at HPV FOCAL trial enrolment and geographical location. Survey respondents and non-respondents were comparable by study arm and marital status, but those who responded to the survey were slightly older than non-responders (median of 51 years vs 49 years), although the difference was not clinically significant.

Table 2 Multivariate analysis of predictors for participants who are accepting of HPV testing instead of Pap testing for cervical cancer screening

Variable	OR (95% CI)	P value
Age at survey completion (years)	1.01 (1.00 to 1.02)	0.01
Received HPV positive results during study (yes vs no)	1.41 (1.11 to 1.80)	0.005
Education (incomplete postsecondary or less vs complete college or higher)	1.06 (0.93 to 1.21)	0.41
Marital status (living without a partner vs living with a partner)	1.04 (0.88 to 1.22)	0.67
Income (\$75 000 or more vs less than \$75 000)	0.97 (0.84 to 1.12)	0.68

HPV, human papillomavirus.

Acceptability of HPV testing for screening

Overall, 63% of survey respondents agreed or strongly agreed that HPV testing for cervical cancer screening was acceptable, with 37% not agreeing (11% disagree, 16% neutral, 10% don't know) (table 1). There were no significant associations between acceptability of HPV testing and marital partnership status, time since study exit, education or income. Women who received an HPV positive result at any point during the HPV FOCAL trial and who were older were more accepting of HPV testing compared with those who remained HPV negative during trial participation.

In multivariate analysis, women who reported HPV testing as acceptable were more likely to have received an HPV positive screen test result at some point during the trial (OR 1.41, 95% CI 1.11 to 1.80, $p=0.005$) and were older (OR 1.01, 95% CI 1.00 to 1.02, $p=0.01$) (table 2).

Over half of respondents (54%) agreed with the statement 'I would be willing to have an HPV test every 4–5 years instead of a Pap every 3 years'. There was a significant difference in acceptability of an extended screening interval between those who reported being accepting of HPV testing compared with those who were not accepting. Overall, 69% responded that HPV testing starting at age 30 years was acceptable, with over 80% of those who were accepting of HPV testing reporting agreement with a higher screening age (30 years or over) compared with Pap testing (table 1).

In addition, 66% of respondents reported that an extended screening interval would not result in less visits to their healthcare provider for other medical reasons, indicating that despite the extended interval recommended with HPV-based screening, women would continue to see their providers for medical reasons as needed.

Attitudes towards an HPV positive test result

Survey respondents were asked to rate their level of agreement with a variety of statements regarding attitudes surrounding receipt of HPV positive results (table 3). Women were asked if they would be more concerned about receiving a positive HPV test or an abnormal Pap result, for which most respondents (73%) reported that both screening outcomes would concern them equally. However, those that reported HPV testing to be acceptable reported that an HPV positive result would concern

them more (14%) than an abnormal Pap. This was statistically different compared with those who were not accepting of HPV testing, who responded that abnormal Pap test results would concern them more.

Most respondents who were not accepting of HPV testing indicated that having a sexually acquired infection would concern them differently than having abnormal Pap results. The relationship between level of agreement with HPV testing acceptability and one's level of concern about having a sexually acquired infection was significant. Regardless of a respondent's reported HPV testing acceptability, most respondents felt it important to them to know who gave them HPV and when they acquired HPV (71% and 78%, respectively). Most respondents indicated they disagreed or were neutral regarding feeling judged for having HPV, and there was no significant relationship between feeling judged and level of agreement with HPV testing. More respondents who were accepting of HPV testing indicated they would feel comfortable telling their partner if they had HPV, which was in contrast to those who were not accepting of HPV testing; these differences indicated a significant relationship between HPV testing acceptance and comfort disclosing HPV status to a partner. Regardless of level of agreement with HPV testing, most respondents (79%) indicated they would feel concerned about transmitting HPV to their partner(s). More respondents who were accepting of HPV testing would feel confident in the recommendations from their provider for the management of their HPV positive results. There was a significant association between level of agreement with HPV testing and degree of confidence with provider recommendations.

Sources of information for HPV testing and screening

Overall, the most reported important sources of information for all respondents were healthcare providers and BC Cancer, the agency that is responsible for the cervix screening programme in British Columbia (table 4). Further, those that are accepting of HPV testing were more likely to look to their healthcare providers and BC Cancer as important sources of information than those not accepting. Regardless of one's level of agreement with HPV testing for screening, friends and family or social media were not as important as healthcare providers and BC Cancer for sources of information.

**Table 3** Experiences receiving HPV positive results

	HPV testing instead of Pap testing for cervical cancer:			P value
	Total (N)	Not acceptable, n (%)	Acceptable, n (%)	
	5336	1993 (36.0)	3343 (60.4)	
What would concern you most: being told you had abnormal Pap results, or being told you had HPV?†				
Being told I have abnormal Pap test results	668	274 (14.0)	394 (11.9)	<0.001*
Being told I have HPV	683	197 (10.0)	486 (14.7)	
Both would concern me equally	3855	1464 (74.7)	2391 (72.2)	
Neither would concern me	67	26 (1.3)	41 (1.2)	
Having an infection that is sexually acquired (HPV) does not concern me any differently than abnormal Pap results would†				
Agree	1412	442 (24.8)	970 (31.5)	<0.001*
Disagree	2465	974 (54.6)	1491 (48.4)	
Neutral	987	369 (20.7)	618 (20.1)	
It would be important for me to know who gave me HPV†				
Agree	3468	1251 (70.3)	2217 (71.7)	0.094
Disagree	545	188 (10.6)	357 (11.5)	
Neutral	859	340 (19.1)	519 (16.8)	
It would be important for me to know when I got HPV†				
Agree	3795	1372 (76.9)	2423 (78.3)	0.131
Disagree	432	151 (8.5)	281 (9.1)	
Neutral	650	260 (14.6)	390 (12.6)	
I think people would judge me for having HPV†				
Agree	1775	663 (37.4)	1112 (36.0)	0.307
Disagree	1419	495 (27.9)	924 (30.0)	
Neutral	1666	617 (34.8)	1049 (34.0)	
I would feel comfortable telling my partner if I had HPV†				
Agree	3391	1198 (67.4)	2193 (71.3)	0.016*
Disagree	709	284 (16.0)	425 (13.8)	
Neutral	755	296 (16.6)	459 (14.9)	
I would not be concerned about transmitting HPV to my partner†				
Agree	515	174 (9.8)	341 (11.1)	0.095
Disagree	3825	1400 (78.7)	2425 (79.0)	
Neutral	509	205 (11.5)	304 (9.9)	
Being HPV positive would not affect my relationship with my partner†				
Agree	1249	445 (25.1)	804 (26.3)	0.035*
Disagree	2003	708 (39.9)	1295 (42.3)	
Neutral	1584	622 (35.0)	962 (31.4)	
Having HPV would not cause me any concern about cervical cancer†				
Agree	181	52 (2.9)	129 (4.2)	0.012*
Disagree	4112	1499 (84.1)	2613 (84.8)	
Neutral	569	231 (13.0)	338 (11.0)	
I would feel confident in the recommendations from my healthcare provider for follow-up of my HPV positive result†				
Agree	3876	1323 (74.0)	2553 (82.5)	<0.001*
Disagree	226	97 (5.4)	129 (4.2)	
Neutral	780	369 (20.6)	411 (13.3)	

*Significant to p<0.05.

†Missing values up to 5336=not reported or prefer not to answer.
HPV, human papillomavirus.

Table 4 Important sources of HPV information

	Total (N)	Having an HPV test instead of a Pap smear to screen for cervical cancer is acceptable to me		
		Not acceptable, n (%)	Acceptable, n (%)	P value
	5336	1993 (36.0)	3343 (60.4)	
Healthcare provider†				
Important	3405	1128 (59.3)	2277 (70.1)	<0.001*
Neutral	1256	560 (29.5)	696 (21.4)	
Not Important	487	213 (11.2)	274 (8.4)	
BC Cancer (organises screening in BC)†				
Important	2713	907 (48.7)	1806 (56.9)	<0.001*
Neutral	1649	673 (36.1)	976 (30.8)	
Not Important	673	283 (15.2)	390 (12.3)	
Government websites (eg, Canadian Cancer Society/Public Health Agency Canada)†				
Important	2111	734 (39.9)	1377 (44.2)	0.011*
Neutral	1914	748 (40.6)	1166 (37.5)	
Not important	928	359 (19.5)	569 (18.3)	
Other websites (WebMd, blogs)†				
Important	1597	567 (31.3)	1030 (33.7)	0.112
Neutral	1997	776 (42.8)	1221 (40.0)	
Not Important	1273	469 (25.9)	804 (26.3)	
Friends/family†				
Important	1336	453 (25.0)	883 (28.9)	0.012*
Neutral	1865	725 (40.0)	1140 (37.3)	
Not Important	1670	634 (35.0)	1036 (33.9)	
Social media†				
Important	580	197 (10.9)	383 (12.6)	0.004*
Neutral	1562	633 (35.1)	929 (30.7)	
Not Important	2693	975 (54.0)	1718 (56.7)	

*Significant to $p < 0.05$.
 †Missing values up to 5336=not reported or prefer not to answer.
 BC, British Columbia; HPV, human papillomavirus.

DISCUSSION

Acceptability of, and attitudes towards primary HPV testing were analysed from 5532 women who completed HPV testing as part of the HPV FOCAL trial, which was embedded within an organised population-based cervical cancer screening programme. Most respondents (63%) were accepting of HPV testing for cervix screening and for screening with HPV testing to begin at age 30 years (69%). Just over half were accepting of HPV testing with the extended screen interval of 4–5 years (54%). Although most women were accepting, the proportion of respondents indicating they disagreed with or were neutral in their acceptance of HPV testing and of extended intervals was higher than we expected considering this was a group of people who were provided with education and information about HPV, HPV testing and cervical cancer. These findings are similar to other studies that indicate women have concerns about the extended interval

recommended with HPV testing,^{10 20–22} stemming from a belief that a cancer diagnosis may be missed through extension of the interval. Considering study participants received information regarding HPV, the natural history of HPV and cervical cancer and the rationale for HPV testing compared with the Pap test, these findings indicate that additional research and patient engagement is needed to gain insights and identify and develop resources or procedures to address barriers to HPV testing and an extended screening interval.

This multivariate analysis found that those who received positive HPV test results at some point during the trial were more likely to be accepting of HPV testing for cervix screening than those who never received an HPV positive result. Those who tested HPV positive would have received additional information and counselling from their healthcare provider and or a study nurse, which would not necessarily have been provided to those who



tested HPV negative. This additional information would have reinforced the education participants were provided at trial baseline, including the prevalence of HPV in the population, the transient nature of HPV and the long natural history between HPV infection and cervical dysplasia development. In addition, those with HPV positive results would have received additional follow-up and management by the time the survey was administered. The reinforcement of education, an opportunity for dialogue when receiving the positive result, and the fact that those with a positive HPV result would have received treatment for detected dysplasia, may have facilitated improvement in knowledge and, subsequently, enhanced acceptance of HPV testing. Other findings have indicated that increased HPV and HPV screening knowledge can be a facilitator of HPV screening acceptance.¹⁸

One of the concerns with an extended screening interval is if women would be less likely to consult with the healthcare provider for other medical reasons.¹⁶ When we asked participants if they would be less willing to see a healthcare provider for other medical reasons if the interval for cervix screening were increased, most respondents said they would not be less willing, indicating that the extended interval for cervix screening would not prevent them from seeking care as needed. This finding can provide reassurance to healthcare professionals who have concerns that the extended screening interval recommended with HPV-based screening would lead to fewer visits to a clinician, given that the cervical screen visit is often an opportunity for the clinician to assess other preventive care or medical issues.²³

Respondents' concerns regarding receipt of HPV positive results were varied. Nearly 93% of the respondents never received an HPV positive result during the HPV FOCAL trial. As a result, the majority of responses evaluate attitudes and experiences for those who did not actually receive HPV positive results, and therefore, reflect how the respondents would hypothetically feel if they were to receive HPV positive results.

Most participants indicated that having HPV would cause them concern about having cervical cancer. The majority of participants reported that having either an abnormal Pap or a positive HPV test would concern them; however, for those that had tested positive for HPV, they reported that an HPV test would concern them more compared with an abnormal Pap. Overall, participants' perceptions about HPV positive results and cervical cancer indicate that increased knowledge regarding the specificity of HPV testing for cervical cancer screening is needed.

Receipt of positive HPV results has been associated with higher anxiety and distress compared with receipt of abnormal Pap results,^{24–26} which may be due to the fact that HPV is an STI and has been associated with levels of shame and stigma.^{27 28} Most respondents in our survey indicated that receiving results for a sexually acquired HPV infection would concern them differently than having an abnormal Pap test result; however, this belief

varied depending on a participant's acceptability of HPV testing. Women who were not accepting of HPV testing indicated that an HPV positive result would concern them differently than abnormal Pap results, compared with those who were accepting of HPV testing. Most respondents, whether they accepted HPV testing or not, felt it important for them to know who gave them HPV and when they got it (71% and 78%, respectively). These findings together are reflective of other research findings^{9 29 30} and indicate that when developing education and communication strategies, emphasis should be placed on the high prevalence of HPV in the population, the transient nature of most HPV infections and bringing awareness to the fact an infection may have been acquired several years prior to a positive test result. Differentiating HPV from other STIs may minimise anxiety and facilitate normalisation and acceptance.^{16 27}

Almost 75% of the respondents indicated they felt an HPV positive result would affect their relationship with their partner, or they were not sure, and almost 80% would be concerned about transmitting HPV to their partners, with many feeling they would be judged for being HPV positive. The stigma associated with HPV, concerns about infidelity and potential partner reactions to the HPV result may underlie these concerns. Previous research has indicated some women question whether partner notification with HPV is necessary.^{16 31} Unlike other STIs such as chlamydia or gonorrhoea where partner notification is recommended for testing and treatment purposes, there is usually no medical reason to notify the partner of a woman who tested positive for HPV.

Healthcare providers, as trusted and valued sources of information, can influence patients' decision-making patterns regarding healthcare decisions.^{20 32} Reflecting other findings, the respondents in this survey indicated that the most important sources of information for them were their healthcare providers and the provincial screening programme.^{20 32} In this cohort, the least important sources of information were social media and friends and family, providing reassurance that women in this cohort seek information from reputable sources such as healthcare providers and the screening programme compared with the internet, or friends and family. These findings demonstrate that programme planning for HPV-based screening should prepare healthcare providers with adequate education and training surrounding HPV prior to programme changes, to ensure they are equipped to address women's questions and concerns regarding the paradigm shift from cytology to HPV-based screening.

This study is not without limitations. Survey participants were part of a large clinical trial and were given information about HPV, HPV testing and cervical cancer on enrolment and, therefore, may not be representative of all people eligible for cervix screening in British Columbia. However, participants of this study are reflective of the current population engaged in the screening programme, who receive cytology testing with the Pap smear and not HPV testing as standard of care. As a

result, their concerns and feedback are informative for programmes planning for a shift from cytology to HPV-based screening. The response rate may be considered low at 42%; however, those who have stronger opinions about their screening choices may have been more likely to respond to the survey than those who are more trusting of the healthcare system and accepting of any future policy changes. In addition, survey respondents were representative of overall participants in the HPV FOCAL trial. The majority of respondents completed the survey approximately 3 years after trial completion, and it is possible there was loss of recall of HPV related information provided to them when they consented to participate in the trial, which for many was up to 7 years prior to survey completion. The potential lag time between trial entry and survey completion may have introduced recall bias and impacted women's attitudes and beliefs surrounding HPV testing; however, the impact of this potential bias would be small as we found no significant difference between acceptability and time between trial entry and survey completion. In addition, most of the participants in this trial were over the age of 50 years, highly educated and primarily from two urban geographic regions and may not be representative of all screen eligible people in various regions of British Columbia.

CONCLUSIONS

In this study within an organised screening setting, evaluating acceptability and attitudes around HPV testing from women undergoing HPV-based screening, most are accepting of HPV testing for screening; however, further research is needed to understand factors that can increase acceptability. These findings contribute to the growing body of evidence demonstrating that concerns and experiences surrounding HPV testing and receipt of HPV positive results are complex and varied. As many cervix screening programmes begin HPV-based screening and are planning implementation strategies, attention to patient engagement to address potential barriers will be important. As HPV-based screening becomes standard of care, it is plausible that concerns with this paradigm shift will eventually be alleviated with increasing knowledge and familiarity. These findings provide insight into areas of importance that should be considered for development of education, information and communication strategies.

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