Randomised controlled trial of the effect of evidence based information on women’s willingness to participate in cervical cancer screening

P Adab, T Marshall, A Rouse, B Randhawa, H Sangha, N Bhangoo

Study objectives: To assess whether providing women with additional information on the pros and cons of screening, compared with information currently offered by the NHS, affects their intention to attend for screening.

Design: Randomised controlled trial. Participants were randomly assigned to receive either the control, (based on an NHS Cervical Screening Programme leaflet currently used), or the intervention leaflet (containing additional information on risks and uncertainties).

Setting: Three general practices in Birmingham.

Participants: 300 women aged 20 to 64 attending the practices during a one month period.

Main outcome measures: Intention to attend for screening.

Main results: 283 women (94.3%) completed the study. Fewer women in the intervention (79%) than the control group (88%) expressed intention to have screening after reading the information leaflet. The crude odds ratio (OR) and 95% confidence intervals (CI) was 0.50 (0.26 to 0.97). After adjusting for other factors, the trend persisted (OR 0.60, 95% CI 0.28 to 1.29). Having a previous Pap smear was the only significant predictor of intention to have screening (adjusted OR 2.54, 95% CI 1.03 to 6.21).

Conclusions: Providing women with evidence based information on the risks, uncertainties, and the benefits of screening, is likely to deter some, but not differentially those at higher risk.

METHODS
Study design and population
This was a randomised controlled trial undertaken at three general practices in Birmingham. We invited women between the ages of 20 and 64 attending the practices to participate. Permission for the trial was obtained from South Birmingham Research Ethics Committee and all participants were provided with information on the study before being asked to give written consent.

Intervention
We devised two types of information leaflet. The control leaflet was based on one produced by the NHSCSP. It included information on the nature and purpose of screening, what the
test involves, choice of venue, how results will be obtained, and possible reasons for further tests.

The intervention leaflet, in addition to the above, contained information on the absolute average individual risk for cervical cancer, likelihood of positive findings, the possibility of false positive/negative results, the uncertainties attached to the screening process, the absolute benefit associated with screening and the cost of the process to the NHS (appendix, available on the journal web site www.jech.com/supplemental).

As the NHS cervical screening programme is well established, and coverage is linked to practice performance measures and to GP remuneration, all reference to “cervical” cancer, “cervical” screening, or “smear test” were removed from the leaflets. This was to gain cooperation and minimise disruption to participating practices. Nevertheless, all the facts presented were related to cervical screening. Thus the intervention leaflet explained that screening prevented the onset of this particular cancer. In addition, it explained the burden of cancer (10/10 000 mortality), the absolute benefit of screening (reduced risk to 1/10 000), risk of false positive (2000 recalled for every one woman with cancer) and false negative (around 10%). This information was all based on data from the cervical screening programme in the UK.

Protocol and random assignment

We developed a structured questionnaire that was previously piloted on 20 women. One of us (PA) prepared a computer generated list of random numbers, which was used to sequence questionnaires to contain either the control or intervention leaflet. Three of us (BR, HS, and NB) visited participating practices on several occasions between April and May 2001 and distributed questionnaires in random order. Participants, who were blind as to which arm they were allocated to, were asked to leave completed questionnaires with the reception before leaving the practice.

Outcome measures

The main outcome measure was expressed willingness to have the “study screening test”. This is an accepted outcome measure, 23 24 having also been used in previous studies.

Other measures

We collected information on sociodemographic factors, health related behaviours, and whether they had any family or close friends with cancer. Social class of respondents was determined based on the occupation of the main wage earner in the household (social class I=highest, and V=lowest). In addition, we developed a risk score for cervical cancer, based on the woman’s age, social class, and smoking status. Those with scores below the median were labelled as “lower risk”, and the rest as “higher risk”.

Sample size

The five year uptake of cervical screening in the UK is estimated to be around 80%. 4 We calculated that in order to detect a difference in intended uptake between groups of 15% or more with 95% confidence and 80% power, a minimal sample size of 276 (138 in each group) was required.

Statistical analysis

We compared intended uptake of the study screening test among the intervention and control groups. Bivariate analysis, using $\chi^2$ for binary categorical variable, $t$ for trend for ordinal variables and $t$ test for linear variables, was used to compare intended screening in relation to other factors. Any characteristic that was associated with intended uptake at a level of significance of 10% or less was entered in a logistic regression model (using the “enter” method), to obtain an adjusted odds ratio for the intervention compared with the control group.

Subgroup analysis was performed to compare women at higher and lower risk of cervical cancer. All statistical analyses were performed with SPSS (version 10).

RESULTS

Participant flow and follow up

About 10% of women approached refused to participate, and were not given a questionnaire. Among 300 women who gave consent, 283 (94.3%) returned their questionnaires (fig 1).

General description

The mean age of responders was 39.4 years, and a sizeable minority (17.3%) were non-white, reflecting the population in Birmingham and the practices participating in the study. Control and intervention groups were similar in all respects, except for a significantly higher proportion of non-white women in the intervention compared with the control group (table 1).

Intended uptake of screening test

Most responders (229 of 274, 83.6%) expressed their willingness to attend for the study screening test. However, those in the intervention group were significantly less likely to want the test (109 of 138, 79.0%) compared with the control group (120 of 136, 88.2%) (difference between groups 9.2% (95% CI 3 to 21.7), unadjusted OR 0.50 (95% CI 0.26 to 0.97) p=0.039).

A high proportion of responders (257 of 277, 92.8%) had attended for a Pap smear in the past; 72.9% within three years, and 82.3% within the past five years. None of the sociodemographic or behavioural factors we inquired about were significantly associated with having had a Pap smear within the past five years. However, women who had been previously screened were significantly more likely to say they would attend for the study screening test (OR 2.3, 95% CI 1.1 to 4.8). Other factors associated with intention to take up the study screening test included attending regularly for dental check ups (OR 3.19, 95% CI 1.52 to 6.73) and having close friends or family with cancer (OR 1.98, 95% CI 0.99 to 3.96). After adjusting for these variables, exposure to the intervention leaflet was still associated with reduced expressed willingness to have the study screening test, though this was no longer statistically significant (table 2). Having had a Pap smear in the past was a significant predictor of intention to have screening, whereas cervical cancer risk was not. Repeating the
logistic regression model with an interaction term between the intervention effect and level of risk showed no significant interaction (p=0.59).

We also found that 87.6% (240 of 274) thought the government should set up a national screening programme, including 59% (n=26) of those who did not want to attend themselves. Those in the intervention group were less likely to think such a programme should be implemented compared with the control group, though the difference was not statistically significant.

**DISCUSSION**

We found that providing women with more information about the risks and uncertainties of screening, as well as the benefits, resulted in a small reduction in expressed willingness to attend for screening. However, even among women who were given more information, intended screening rates were nearly 80%.

**Strengths and weaknesses of the study**

To our knowledge this is the first trial to assess the effect of giving evidence based information on women’s expressed willingness to attend for a screening test that is already well established. The five year coverage for cervical screening within the study population was the same as that for Birmingham generally (80.6%), and this, together with the high response rate suggests that they were fairly representative of the target group.

We tried to blind participants to what the study test was, and the condition to which screening referred. The aim was to limit interference with their prior understanding and beliefs about the Pap smear, and to gain cooperation from participating practices. On the other hand, decisions may have differed had women known that we were referring to cervical screening. As in some other studies, our principal outcome was expressed willingness to have screening, which may differ from actual attendance. While actual uptake would be the ideal outcome measure, time and resource limitations prevented us from using this. This may partly explain the higher proportion of women who said they would attend in the control group, compared with those that currently have Pap smears. We could not compare knowledge between the two groups, and cannot directly infer that women in the intervention group had a better understanding of the pros and cons of the test. However, evidence suggests that people generally understand absolute risks (as used in the intervention leaflet) better than relative risks. Also, our sample size did not allow sufficient power to detect a difference in intended uptake of less than 15% between groups.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Comparison of intervention and control groups at baseline (values are numbers and (percentages) unless otherwise stated)</th>
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<tbody>
<tr>
<td>Participant characteristics</td>
<td>Control (n=141)</td>
</tr>
<tr>
<td>Sociodemographic characteristics</td>
<td></td>
</tr>
<tr>
<td>Age*</td>
<td>39.3 [20–64]</td>
</tr>
<tr>
<td>Marital status</td>
<td></td>
</tr>
<tr>
<td>Single/divorced/widowed</td>
<td>54 [39.1]</td>
</tr>
<tr>
<td>Married/living with partner</td>
<td>84 [60.9]</td>
</tr>
<tr>
<td>Social class</td>
<td></td>
</tr>
<tr>
<td>I and II</td>
<td>35 [24.9]</td>
</tr>
<tr>
<td>III</td>
<td>58 [41.3]</td>
</tr>
<tr>
<td>Ethnicity</td>
<td></td>
</tr>
<tr>
<td>White</td>
<td>118 [84.1]</td>
</tr>
<tr>
<td>Non-white</td>
<td>17 [12.6]</td>
</tr>
<tr>
<td>Personal contact with someone with cancer</td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>64 [45.4]</td>
</tr>
<tr>
<td>Yes</td>
<td>77 [54.6]</td>
</tr>
<tr>
<td>Having family or close friends with cancer</td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>28 [20.3]</td>
</tr>
<tr>
<td>Yes</td>
<td>120 [85.7]</td>
</tr>
<tr>
<td>Smoking status</td>
<td></td>
</tr>
<tr>
<td>Never smoker</td>
<td>68 [48.2]</td>
</tr>
<tr>
<td>Former smoker</td>
<td>36 [25.5]</td>
</tr>
<tr>
<td>Current smoker</td>
<td>37 [26.2]</td>
</tr>
<tr>
<td>Risk for cervical cancer</td>
<td></td>
</tr>
<tr>
<td>Risk score* (based on age, social class, and smoking status)</td>
<td>5.5 [3–9]</td>
</tr>
</tbody>
</table>

*Values are mean (range).

<table>
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<tr>
<th>Table 2</th>
<th>Results of logistic regression model assessing the factors associated with women’s expressed intention to have the study screening test</th>
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</thead>
<tbody>
<tr>
<td>Intention to have study screening test</td>
<td>Adjusted OR (95% CI)</td>
</tr>
<tr>
<td>Exposure to intervention</td>
<td>0.60 (0.28 to 1.29)</td>
</tr>
<tr>
<td>Previous Pap smear within past five years</td>
<td>2.54 (1.03 to 6.21)</td>
</tr>
<tr>
<td>Regular dentist attendee</td>
<td>2.26 (0.96 to 5.29)</td>
</tr>
<tr>
<td>Having family/friends with cancer</td>
<td>1.99 (0.89 to 4.48)</td>
</tr>
<tr>
<td>Higher risk for cervical cancer</td>
<td>0.94 (0.44 to 2.02)</td>
</tr>
</tbody>
</table>
Key points

- Cervical screening can cause harm as well as benefit.
- Despite guidance from the GMC and others to obtain informed consent before screening, the risks and uncertainties were not well communicated to women.
- There are fears that giving women additional evidence based information on screening will reduce uptake and undermine the national programme.
- We found that providing additional, more balanced information about screening does reduce willingness to attend for screening, but not considerably so.
- The findings should counteract the reluctance among some to provide women with more information.

Policy implications

- Ethical and professional considerations as well as social demand are all drivers for involving the public in decisions about their participation in screening.
- Our findings suggest that providing women with more balanced information on the risks as well as the benefits of screening would not undermine the national screening programme.
- Given the paucity of relevant trials, our findings are important in triggering further research. Further studies are needed to confirm our findings, using actual attendance as an outcome.

Findings in relation to other studies

A few studies have investigated the effects of offering different types of information on intended or actual uptake of screening. Providing more information increases knowledge and assists in decision making, but has an unpredictable effect on uptake. In three studies assessing the effect of information giving on decisions to undergo screening for prostate cancer, two found that intervention reduced uptake, while in the other there was no effect. In a study to assess willingness to undergo screening for pancreatic cancer, participants who were given extended information were significantly less likely to accept the test compared with those given basic information. However, in a trial of women at low to moderate risk of breast cancer, better information had no effect on women's willingness to have genetic screening. Decisions on screening are not just influenced by the information provided, but also by other factors, such as values, cultural beliefs, and personal experiences. Indeed, we found a tendency for women who had personal contact with someone with cancer to be more likely to want to undergo screening.

Unanswered questions and future research

Our intervention leaflet offered evidence based information related to the topics emphasised by the GMC. However, we do not know whether this contained sufficient information for decision making and whether it included messages, that women who have been through the process themselves, would feel are important. Furthermore, although there is some evidence that the medium used to convey information has little effect on knowledge, understanding, or decision making, little is known about the most effective form of presentation.

Conclusions

Our findings suggest that providing women with a more balanced appraisal of the pros and cons of screening, as well as being more ethical, would result in a reduction in screening uptake, though our point estimate suggests that uptake rates would not be much below the current target rate of 80%. Furthermore, we found no evidence that information would adversely affect women at higher risk.

ACKNOWLEDGEMENTS

We thank all the staff at Colton Medical Centre (Longbridge), Jiggin’s Lane Practice (Bartley Green), and Shanklin House Surgery (Handsworth) for their cooperation during this study. We also thank Dr Paul Aveyard for his help with some of the analysis.

Contributors

PA conceived the idea for the project, supervised it, and wrote the final paper. BR, HS, and NB developed the project, carried out the data collection, collation and initial analysis, and wrote the first draft. TM and AR were involved in initial discussions on the project and in developing the hypothesis, calculated the population effects of cervical screening, and provided this for the information leaflets used, and have contributed to the final draft.

A copy of the intervention leaflet used in the study is available on the journal website (www.jech.com/supplemental).

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Health, equity, human rights, and the invasion of Iraq

During the days culminating in the US-British invasion of Iraq, I was working with a human rights attorney colleague, exploring links and distinctions between health, equity, and human rights. These concepts thus have been in my thoughts, and concern for each leads me to deplore the invasion of Iraq on several counts.

Firstly, given the scale of suffering and death that inevitably accompany war, it is unconscionable to embark on that course, even for a just cause, except as a clearly demonstrated last resort. There is widespread international consensus that reasonable alternatives to war had not been exhausted in this case. The consequences of war include not only its direct effects but the massive aftershocks resulting from destruction of infrastructure (for example, clean water) critical for survival and health; many more deaths occurred for this reason in the wake of the first Gulf War than as a direct result of the military action itself. Moreover, this is not a just war; evidence linking Iraq and September 11 or Al Qaeda was never produced, and equally heinous regimes have been tolerated or supported (as Saddam Hussein was previously) by the US. There are many reasons to suspect that the real motives for US interest in a regime change in Iraq have more to do with control of oil and empire than with fighting terrorism. Terrorism will surely increase in light of the hatred this war and the ensuing occupation will provoke for generations to come, throughout the Arab and Islamic worlds and among others who reject the disturbing vision of a hegemonic New World Order evoked by this invasion.

Secondly, by setting the frightening dual precedents of preemptive military strikes and defiance of the United Nations, this action drags the entire world backward toward the laws of the jungle, obliterating decades of work toward global disarmament and 50 to 100 years of work toward international governance. The arrogance this reflects is in itself shocking.

Thirdly, this war will exacerbate inequities. In Iraq, it undoubtedly is now taking and will continue in its aftermath to take its heaviest toll on the poor and especially poor children. In the US, the costs of the war and its aftermath most certainly will accelerate the dismantling of public services already started by the current administration's domestic policies, thereby increasing social disparities in this country; and there will be less support for international development outside the self serving agenda for the Fertile Crescent.

Fourthly and finally, this action represents a grave threat to human rights globally. By its explicit undermining of the authority of the United Nations (UN), the Bush administration has implicitly undermined the force of international law overall and specifically of human rights treaties and other agreements developed under UN auspices. We must publicly condemn this unjustified war, find ways to help repair the damage, and develop new strategies to struggle for health, equity, and human rights in a world that is far more brutal and violent than the world we had dared hope to encounter in the 21st century.

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