Facilitating chlamydia testing among young people: a randomised controlled trial in cyberspace

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ABSTRACT

Objectives Chlamydia notifications have been rising in Australia for over a decade and are highest in young people. This study aimed to evaluate the impact of an internet-based intervention on chlamydia testing among young people 16–25 years.

Methods In this randomised controlled trial, recruitment, data collection, study interventions and follow-up occurred entirely in cyberspace, facilitated by a website. Eligible participants were aged 16–25 years and resided in Australia. The intervention group received personalised emails inviting interaction about chlamydia testing, while the control group received regular impersonal emails. Primary outcome was self-reported chlamydia testing at 6-month follow-up; secondary outcomes were condom use and changes in knowledge and attitudes.

Results 704 young people completed baseline information; 40 were excluded and five withdrew prior to follow-up. The follow-up rate was 47.3% overall. In the intervention group, 40.6% (95% CI 30.7% to 51.1%) reported having had a chlamydia test at follow-up compared with 31.0% (95% CI 24.8% to 37.2%) in the control group (p = 0.07). A per-protocol analysis found that those who engaged in email interaction were more likely to report chlamydia test uptake compared with those in the control group (52.5%, 95% CI 39.3 to 65.4% cf 31.0%, 95% CI 24.8% to 37.2%, p = 0.002). There were no differences in secondary outcomes between groups.

Conclusions This is the first randomised controlled trial undertaken in cyberspace to promote chlamydia testing. E-technology may be useful in promoting chlamydia testing and healthcare seeking behaviour in young people.

INTRODUCTION

Among the sexually transmissible infections (STIs) other than HIV, chlamydia causes the greatest burden of disease globally.1 In Australia, notification rates for genital chlamydia infection have steadily increased over the past decade and are highest in women and men aged 15–24 years.2 In 2011, notification rates were 1443.3 per 100 000 among 15–19-year-olds and 1901.4 per 100 000 among 20–24-year-olds.2 Australia’s first National STI strategy 2005–2008 identified young people aged 16–25 years as a target group for chlamydia control and prevention.3 This study was one of several pilot projects to inform a national screening programme3 and aimed to evaluate an internet-based intervention to increase chlamydia testing.

Australian young people experience several barriers to care, including lack of knowledge about services available and how to access them, concerns about confidentiality, embarrassment, cost and transport.5 However, most young people are familiar with and access general practice.6 An analysis of general practice encounters across Australia from 2000 to 2007 found that the rate of chlamydia testing among 15–24-year-old patients was only 13.2 per 1000 encounters.7 Chlamydia testing rates among young people are higher in sexual health and family planning clinics,8 but unlike general practices, these are not located in many rural and some urban areas. Thus, the majority of the youth population who might be at risk are unlikely to be tested for chlamydia.

Over 85% of Australian young people access the internet each year9 and young people find it useful for seeking help around sensitive issues. Furthermore, men are as likely as women to use the internet to seek help.9–11 A study in Switzerland suggested that the internet can facilitate access to healthcare through the use of tailored information provided to individuals in response to questions asked via a website.12

This study used the internet to engage sexually active young people aged 16–25 years residing in Australia, in confidential personalised email interactions with a clinician, with the goal of facilitating their access to primary health services for chlamydia testing. The primary aim of this study was to evaluate the impact of this intervention on chlamydia testing when compared with a control intervention (monthly impersonal emails for 6 months).

METHODS

Study location and population

This was a randomised controlled trial (RCT) with 1:2 randomisation. Eligible participants were aged 16–25 years residing in Australia who had had penetrative sexual intercourse and who provided a valid email address. Eligibility was determined by self-report of these criteria.

The study setting was cyberspace. A website (http://www.getcluedup.com.au) was the vehicle for accessing the intervention and was developed in consultation with 20 youth consultants (16–25 years) who were recruited through professional and collegiate networks. Website content included information about chlamydia and testing but also addressed known gaps in knowledge13 and barriers to seeking help14 and provided service.
directories and links (see web appendix 1). The website went live in March 2007 with recruitment into the study commencing simultaneously. Recruitment ceased in January 2008, but the website remains live. The website was promoted via paid advertising, existing youth websites, social networking sites and opportunistic media interviews. Google Analytics was used to monitor website traffic.

The website invited eligible visitors to participate in the study via a homepage banner and clickable tiles on the other pages. These links took visitors to the participant information statement. Potential participants then entered a current email address and ticked a consent box. This third step took them to the baseline questionnaire housed within the website. The baseline questionnaire collected data on demographics, education/occupation, substance use, sexual history, previous STIs, knowledge about chlamydia and attitudes to chlamydia testing. Participants could go into a draw for an iPod® or a music store voucher if they completed all elements of the study.

**Interventions**

The intervention group received personalised emails from a clinician (sexual health nurse or doctor). A ‘personalised email’ was sent from the clinician’s mailbox, included the clinician’s name and position, and contained a link to their staff profile on the University of Sydney’s website. The email thanked the young person for their participation and said that the clinician would like to ‘chat about chlamydia and getting tested’. The participant was invited to ask questions and prompted with questions about testing knowledge. Young people who responded were then engaged appropriately: advice depended on the questions asked. Non-responders were sent weekly emails for 3 months and then monthly emails for another 3 months. All email communication to non-responders in this group remained personalised, as described above.

Participants assigned to the control group received an email sent from the project mailbox (‘Clued Up’), was signed ‘The Clued Up Research Team’ and did not mention a clinician by name. These emails thanked the young person for participation and stated that they would be sent a reminder email about their participation in the study every month for 5 months and a final questionnaire in 6 months. These emails were intended to enhance retention and completion of the final questionnaire but were not personalised. There was no interaction and no clinical advice provided.

Web appendix 2 gives examples of email interactions.

**Outcomes**

Follow-up took place 6 months after enrolment. Data were collected via an online questionnaire using similar questions to baseline on knowledge, attitudes, sexual history and chlamydia testing in the past 6 months. This questionnaire was accessed via a link sent by email from the clinician. Weekly email reminders were sent for 3 months to non-responders.

The primary outcome was self-report of having had a chlamydia test within the past 6 months. Secondary outcomes were changes in knowledge about chlamydia, attitudes towards chlamydia testing and frequency of condom use.

**Sample size**

A minimum of 320 participants in each group was required to detect at least an 8% positive difference in testing rates in the intervention versus control group with a power of 90% and significance taken at 0.05. A loss to follow-up of 10% in the intervention group and 50% in the control group was anticipated, giving a target sample size of 1000 (360 intervention and 640 control).

**Randomisation**

Allocation to intervention or control groups was done on a 1:2 ratio based on the anticipated loss to follow-up described above. For allocation, a computer-generated random number (1, 2 or 3) was obtained by the same clinicians who sent emails to study participants. The number ‘1’ allocated the young person to the intervention group and ‘2’ or ‘3’ to the control group. Allocation took place each time a young person was deemed eligible (downloaded data showed email address, eligible age, an Australian postcode and ticked ‘Yes’ to having had intercourse) without any other information about the participant being known.

**Analysis and statistical methods**

Both intention-to-treat (ITT) and per-protocol (PP) analyses were performed. Only a proportion of young people in the intervention group interacted with the clinician as per the study protocol, that is, engaged. Engagement was defined as having had a minimum of one response of any type from the young person within 3 months from enrolment. The intervention was only deliverable when engagement occurred, where it did not occur, there was a ‘failure to start the intervention’. Thus, comparisons between the engaged group and the control group constituted the PP analysis. We propose that the PP analysis provides an ‘explanatory investigation of efficacy’ and provides useful information.

Statistical analysis was performed with SPSS V16 with the individual as the unit of analysis. Proportions are presented with 95% CIs. To assess differences at baseline between the intervention (all), engaged and control groups, independent samples’ t tests for the continuous variables and χ² tests for the categorical variables were performed. χ² Analysis was used to assess the statistical significance of differences in the primary outcome and in condom use between groups at follow-up. To adjust follow-up values for baseline values, analysis of covariance was used for knowledge questions and logistic regression for binary outcome measures.

The study was approved by the University of Sydney Human Research Ethics Committee and enrolled in the Australian New Zealand Clinical Trials Registry (ACTRN1260700052459).

**RESULTS**

**Website traffic**

Reliable data on website traffic for the first 3 months (March to May 2007) were unavailable due to technical problems. Traffic between June 2007 and January 2008 ranged from 2030 to 3584 unique visitors per month (see web appendix 3).

**Sample**

Seven hundred and four young people completed baseline information between March 2007 and January 2008 and were randomised. Recruitment was then stopped because of project timelines and budget constraints. Forty young people were subsequently excluded because their email addresses were invalid (all sent emails bounced). Five withdrew prior to follow-up. Three hundred and forty-seven participants did not respond to the 6-month follow-up request. Thus, 312 were included in the ITT analysis (see figure 1).

We compared baseline information between intervention (all), engaged and control groups. Of the baseline sample of 664...
young people, 78.2% were were. The mean age of female participants (20.0 years) was significantly lower than the mean age of male participants (21.5 years), p<0.0001. There was a small significant difference between the engaged and whole intervention groups with respect to being born overseas. Those who engaged were more likely to have been born in Australia. However, of those born overseas in both groups, most were born in English-speaking countries. Table 1 describes their demographic and sexual history characteristics. At baseline, 111/664 (16.7%, 95% CI 13.9% to 19.6%) reported having ever had chlamydia.

The follow-up rate was 47.3% (312/659) overall. There were no differences in demographic (mean age, sex) or baseline sexual history characteristics (number of sexual partners ever, condom use) between those who completed follow-up questionnaires and those who did not (data not shown). There was no difference in follow-up rate between the intervention (49.5%; 96/194) and control (46.5%; 216/465) groups; however, follow-up rate for the ‘engaged’ group (78.2%; 61/78) was significantly higher than for the ‘non-engaged’ group (30.7%; 35/114), p<0.0001, and for the control group (46.6%, 216/464), p<0.0001.

Table 2 presents the primary outcome measurement in the intervention and control groups (ITT) and in the engaged and control groups (PP analysis). Given the loss to follow-up at 6 months is a problem for ITT analysis, we also conducted a sensitivity analysis to examine the possible effects. A higher proportion of young people in the intervention group reported a chlamydia test compared with the control group (40.6% vs 31.0%) but this was not significant. However, the difference between the engaged intervention and control groups was significant (52.5% and 31.0%, respectively, p=0.002).

Of those who reported having had a chlamydia test at follow-up, a total of 14/99 (14.1%, 95% CI 7.95% to 22.6%) young people reported their tests were positive for chlamydia. Three were from the engaged group and 11 from the control group.

The proportion of young people who reported using condoms every time they had sex increased in the engaged and the control groups; however, this increase was not significant. At baseline, 12/61 (19.7%) in the engaged group and 27/216 (12.5%) in the control group reported using condoms always, at follow-up these proportions were 20/61 (32.8%) and 59/216 (27.3%), respectively (p=0.30).

There was no change in knowledge between baseline and follow-up; however, baseline knowledge was high. The proportion of young people in total (n=312) who answered each of the seven questions correctly ranged from 77.6% to 95.8% at
questions and attitude statements are shown in web appendix 4.

Engaged and control groups at follow-up. The list of knowledge differences between the intervention and control or groups or in the engaged and control groups. There were no between change in knowledge in the intervention and control baseline and 80.8 to 97.4% at follow-up. There was no difference in knowledge.

Table 1 Baseline demographic information, sexual history and substance use

<table>
<thead>
<tr>
<th></th>
<th>Intervention (all), n = 196</th>
<th>Intervention (engaged), n = 79</th>
<th>Control, n = 468</th>
<th>Intervention (all) versus control, p Value</th>
<th>Intervention all versus control, p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean age (years)</td>
<td>20.5</td>
<td>20.7</td>
<td>20.3</td>
<td>0.50</td>
<td>0.42</td>
</tr>
<tr>
<td>Female (%)</td>
<td>154 (78.6)</td>
<td>65 (82.3)</td>
<td>365 (78.0)</td>
<td>0.46</td>
<td>0.97</td>
</tr>
<tr>
<td>Not born in Australia (%)</td>
<td>21 (10.7)</td>
<td>4 (5.1)</td>
<td>60 (12.8)</td>
<td>0.06</td>
<td>0.53</td>
</tr>
<tr>
<td>Aboriginal/Torres Strait Islander (%)</td>
<td>4 (2.0)</td>
<td>2 (2.5)</td>
<td>14 (3.0)</td>
<td>1.00</td>
<td>0.70</td>
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<tr>
<td>Region of residence</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Major city (%)</td>
<td>147 (75.0)</td>
<td>54 (68.4)</td>
<td>317 (67.7)</td>
<td></td>
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</tr>
<tr>
<td>Inner regional (%)</td>
<td>33 (16.8)</td>
<td>17 (21.5)</td>
<td>88 (18.8)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Outer regional (%)</td>
<td>12 (6.1)</td>
<td>5 (6.3)</td>
<td>43 (9.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Remote (%)</td>
<td>0 (0.0)</td>
<td>0 (0.0)</td>
<td>4 (0.9)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Very remote (%)</td>
<td>0 (0.0)</td>
<td>0 (0.0)</td>
<td>0 (0.0)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>School (%)</td>
<td>35 (17.9)</td>
<td>11 (13.9)</td>
<td>98 (20.9)</td>
<td></td>
<td>0.70</td>
</tr>
<tr>
<td>University/tertiary (%)</td>
<td>60 (30.6)</td>
<td>26 (32.9)</td>
<td>128 (27.4)</td>
<td></td>
<td>0.42</td>
</tr>
<tr>
<td>Full time work (%)</td>
<td>62 (31.6)</td>
<td>27 (34.2)</td>
<td>149 (31.8)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Part time or casual work (%)</td>
<td>23 (11.7)</td>
<td>7 (8.8)</td>
<td>37 (7.9)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Looking for work (%)</td>
<td>8 (40.8)</td>
<td>3 (3.8)</td>
<td>31 (6.6)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Work + study (%)</td>
<td>4 (2.0)</td>
<td>3 (3.8)</td>
<td>4 (0.9)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Parenting/domestic (%)</td>
<td>2 (1.0)</td>
<td>1 (1.3)</td>
<td>12 (2.6)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other (%)</td>
<td>2 (1.0)</td>
<td>1 (1.3)</td>
<td>7 (1.5)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean age of first intercourse (years)</td>
<td>16.2</td>
<td>16.5</td>
<td>16.3</td>
<td>0.15</td>
<td>0.42</td>
</tr>
<tr>
<td>Mean no. of sexual partners ever</td>
<td>3.1</td>
<td>3.2</td>
<td>3.1</td>
<td>0.83</td>
<td>0.57</td>
</tr>
<tr>
<td>Mean no. of sexual partners last 12 months</td>
<td>2.0</td>
<td>1.9</td>
<td>2.0</td>
<td>0.31</td>
<td>0.88</td>
</tr>
<tr>
<td>Use condoms always (%)</td>
<td>41 (20.9)</td>
<td>20 (25.3)</td>
<td>100 (21.4)</td>
<td>0.62</td>
<td>0.95</td>
</tr>
<tr>
<td>Had chlamydia test in past 6 months (%)</td>
<td>60 (30.6)</td>
<td>26 (32.9)</td>
<td>127 (27.1)</td>
<td>0.49</td>
<td>0.57</td>
</tr>
<tr>
<td>Had a previous diagnosis of chlamydia (%)</td>
<td>26 (13.3)</td>
<td>13 (16.5)</td>
<td>85 (18.2)</td>
<td>0.56</td>
<td>0.21</td>
</tr>
<tr>
<td>Substance use history</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Daily smoker past 12 months (%)</td>
<td>37 (18.9)</td>
<td>17 (21.5)</td>
<td>102 (21.8)</td>
<td>0.80</td>
<td>0.44</td>
</tr>
<tr>
<td>Drink 5 or more standard drinks at a time (%) past 2 weeks</td>
<td>93 (47.4)</td>
<td>47 (59.5)</td>
<td>201 (42.9)</td>
<td>0.67</td>
<td>0.95</td>
</tr>
<tr>
<td>Marijuana use (any) past month (%)</td>
<td>30 (15.3)</td>
<td>10 (13.9)</td>
<td>81 (17.3)</td>
<td>0.45</td>
<td>0.66</td>
</tr>
<tr>
<td>Other illicit drug use (any) past month (%)</td>
<td>23 (11.7)</td>
<td>10 (13.9)</td>
<td>52 (11.1)</td>
<td>0.60</td>
<td>0.83</td>
</tr>
</tbody>
</table>

Baseline and 80.8 to 97.4% at follow-up. There was no difference between change in knowledge in the intervention and control groups or in the engaged and control groups. There were no attitude differences between the intervention and control or engaged and control groups at follow-up. The list of knowledge questions and attitude statements are shown in web appendix 4.

The amount of clinician time required to respond to emails for participants who engaged in interaction was low (a few minutes per email). Responses usually provided information about what testing involved, confidentiality and services available. Some young people asked about other screening tests such as Pap smears, and some were interested in other STIs.

DISCUSSION

To our knowledge, this is the first RCT of an intervention to increase chlamydia testing implemented entirely in cyberspace. Young people who engaged in email interaction with a clinician were more likely to report having a chlamydia test after 6 months than young people who were not offered such contact. Knowledge, attitudes and condom use did not change.

Strengths of this study include originality, youth consultant advice and experimental rigour despite the uncontrollable study environment. Clinicians provided clear information about testing, treatment and relevant services rather than messages about ‘risk’ that could provoke fear. They answered young people’s questions regardless of topic, thus feeling listened-to might have generated confidence in seeking healthcare.

There were several limitations. Website reach was modest and target sample size was not achieved due to project constraints. This combined with loss to follow-up reduced the power of the study. The main outcome measure was based on self-report of testing that could not be verified. Overall follow-up rate was just under 50%; however, of those who engaged, 61/78 (78.2%) completed follow-up. Of the intervention group,

59.4% did not engage. Although there were no measurable differences in attitudes between these subgroups, it is possible that those who engaged were more concerned about chlamydia, more engaged with the health system or wanted to please the clinician or avoid embarrassment. Internet-based trials have high attrition rates, especially when there is no clinician contact at the outset. Attrition can occur either at uptake of an intervention or with failure to complete follow-up. This introduces dilemmas with analysing ITT populations in exactly the way we found. To improve validity, it has been suggested that individuals who do not take up an intervention be removed and that the remaining participants in the intervention group undergo a second randomisation process. Given our time frame, this was not feasible. Most internet-based RCTs have evaluated therapeutic interventions, such as pain management, chronic illness or mental health treatments. It is possible that participants who are asymptomatic are more likely to take up an intervention but also to dropout if no benefit is perceived. Our trial involved an intervention that was not therapeutic and targeted a behaviour change for an asymptomatic condition. We might therefore expect a higher dropout rate compared with therapeutic interventions but this was not the case. Nevertheless, we must be cautious about how to interpret our findings.

Other studies have used e-technology to increase chlamydia testing among young people. A RCT using SMS and email delivered sexual health messages to 994 young people who were recruited from a music festival. The study found that at 12 month follow-up, young women in the intervention group were significantly more likely to report Chlamydia testing than those in the control group. Follow-up rates were substantially lower than ours (34%).

Strategies combining the internet with home-based chlamydia testing have been evaluated. In Sweden, 62.3% of kits requested over the internet were returned for testing,20 in the USA 32.4% of women and 51% of men returned samples and in the Netherlands, 20% of the invites requested testing kits and had returned samples.21 Although our sample size was small, the fact that over 50% of young people who engaged in email interaction reported chlamydia testing at follow-up without the convenience of a home testing kit suggests that email interaction could help facilitate access to health services more broadly.

We found no significant changes in knowledge, attitudes or condom use. However, baseline knowledge was high and might have been positively impacted simply from navigating the website. That reported condom use was low and did not increase significantly highlights the complexities of this behaviour and relationship dynamics at the point of sexual encounter. Further, heterosexual couples using other contraceptive methods are less likely to use condoms or pregnancy might be unexpected. Women are more likely to access chlamydia testing than those who did not engage.

Chlamydia is a common curable sexually transmitted infection that can have potentially serious consequences if untreated and untreated. E-technology has the potential to be a useful adjunct to a population-based screening programme.

Conducting studies in cyberspace is novel and presents challenges to the conventional frameworks for evaluating and interpreting scientific data.

The internet can be a useful vehicle for sexual health promotion and an adjunct to screening strategies.

Key messages
- Young people who engaged with clinicians online were more likely to access chlamydia testing than those who did not engage.
- Conducting studies in cyberspace is novel and presents challenges to the conventional frameworks for evaluating and interpreting scientific data.
- The internet can be a useful vehicle for sexual health promotion and an adjunct to screening strategies.

References

Approval Ethics approval Ethics approval was provided by the University of Sydney Human Research Ethics Committee.

Provenance and peer review Not commissioned; externally peer reviewed.