“Your risk is low, because …”: argument-driven online genetic counselling

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Advances in genetic research have created the need to inform consumers. Yet, the communication of hereditary risk and of the options for how to deal with it is a difficult task. Due to the abstract nature of genetics, people tend to overestimate or underestimate their risk. This paper addresses the issue of how to communicate risk information on hereditary breast and ovarian cancer through an online application. The core of the paper illustrates the design of OPERA, a risk assessment instrument that applies the UK National Institute of Health and Clinical Excellence’s guidelines on the basis of (i) the number of relatives on the same side of the family with the same cancer or cancers that are known to run together; (ii) the ages of these relatives at diagnosis and (iii) the closeness of the family relationship with the person who is doing the assessment. By relying on the argumentation theory, we explain how the communication strategy that OPERA implements is essentially based on Perelman and Olbrechts-Tyteca’s deductive argumentation by association. By using as premises “facts” (propositions about reality that can be assumed without further justification) and “truths” (propositions that make connections about facts), OPERA delivers its claims with an *ex auctoritate* causal link aimed at transferring the audience’s acceptance of the cause to the effect. Overall, the design of OPERA rests on its capacity to induce users’ active processing of risk information through an appeal to their reasoning faculty. In the conclusion, we present some results from a pilot evaluation of users’ acceptance of OPERA.

**Keywords:** internet-enhanced health communication; risk communication; risk-assessment tools; genetic counselling; hereditary breast and ovarian cancer; Toulmin; New Rhetoric; appeal to reasonableness

1. Introduction

Cancer-backup is the leading provider of information for cancer patients in the UK. On January 2008, it merged with MacMillian Cancer Support. On 22 April 2008, Cancer-backup launched Online Patient Education and Risk Assessment (OPERA). OPERA is an online interactive software program – designed by the Institute of Communication and Health and the Technology-Enhanced Communication Laboratory of the University of Lugano (Switzerland) – that offers personalised information about the risk of having an inherited genetic link for breast and/or ovarian cancer in one’s own family. Indeed, 10% of all breast cancer cases have a strong hereditary component relating to mutations on the genes BRCA1 and BRCA2. In addition, women carrying these mutations are at significant risk to develop ovarian cancer (De Grève, Sermijn, De Brakeleer, Ren, and Teugels 2008). OPERA is currently available at: http://www.cancerbackup.org.uk/Aboutcancer/Genetics/GeneticBreastOvarianCancerRiskAssessmentTool.

The communication of risk information on hereditary breast cancer is fraught with difficulties that people have in understanding and evaluating information on risks (Bennet and Calman 1999).
Knowledge regarding risk factors for hereditary breast cancer and the availability of specific options have been shown to be limited or partial (Bottorff, Ratner, Johnson, Lovato, and Joab 1998; Bosompra et al. 2000; Honda and Neugut 2004; Kwate, Thompson, Valdimarsdottir, and Bjoøvberg 2005; Wold, Byers, Crane, and Ahnen 2005; O’Doherty and Suthers 2007). Also people’s general perception of their level of risk can be mistaken. The majority of women who have a relative with breast cancer are not at significantly increased risk. Cancer is, however, an emotive word within the general population. Breast cancer has a particularly high political and media profile, and there appears to be a real, but probably exaggerated fear of developing breast cancer in family members of women who have already developed the disease (Bottorff et al. 1998; Hallowell and Murton 1998; Croyle and Lerman 1999; Rees, Fry, and Cull 2001; Hopwood, Shenton, Laloo, Evans, and Howell 2001). Part of this might stem from the fact that the real extent of a risk translates into assessments, decisions and actions that strongly depend on people’s subjective considerations (O’Doherty and Suthers 2007).

As a matter of fact, the way a person understands his/her risk of carrying a BRCA1 or BRCA2 mutation can negatively impact the actions subsequently undertaken. People who have a significant history of breast and ovarian cancer in the family are offered the genetic test by the National Health Service. Here, it is important that these get to know their breast cancer predisposition and take action accordingly (De Grève et al. 2008). Those who have a negative result for disease-conferring mutations might engage in screening programmes only at a level recommended for the general population. But private genetic testing is available. Considering that genetic testing is an expensive, time-consuming and emotionally demanding procedure, it is crucial that people correctly evaluate whether or not to ask for the private option.

With all this in mind, the central goal of OPERA is to help people reach or shape a realistic understanding of their level of risk and the available options. This paper explains the design of OPERA with a focus on the communication challenges encountered during its development. The material discussed here is the outcome of an initial idea of argumentation-driven online genetic counselling that we presented a few years ago in Mackay, Schulz, Rubinelli, and Pithers (2007). In the concluding section of the paper, we suggest possible strengths in OPERA’s interactional strategy by relying on a set of data from a pilot evaluation of the instrument.

2. The theoretical framework: risk explanations as arguments

The choice of the message delivered to users by OPERA was a particularly critical one to make. The way a message is structured highlights some features of an issue and makes them salient (i.e. more meaningful and memorable) to an audience (Entman 1993).

In the last few years, due to an increased availability of genetic tests and a growth in genetic counselling, the study of risk communication in genetics – especially in the field of hereditary breast and ovarian cancer – has produced several important contributions (Lerman et al. 1995; Lipkus, Klein, and Rimer 2001; Huiart et al. 2002; Skinner et al. 2002; Contegiacomo et al. 2004; Lobb et al. 2004, 2006; Phelps, Platt, France, Gray, and Iredale 2004; Fransen, Meertens, and Schrander-Stumpel 2006). So far, however, there is no consensus on the value of specific strategies of communication on users’ cognitive, affective and behavioural outcomes together (Edwards et al. 2006, 2008). What counts as a successful strategy of risk communication is even less clear in dealing with online instruments such as OPERA, which have the potential of reaching vast audience, but can be affected by the limitations of the electronic channel (Souka and Yoshida 2005). Thus, Green et al. (2004), showed that an online system was more effective than traditional counselling for increasing knowledge of breast cancer and genetic testing. Yet, traditional counselling was more effective at reducing women’s anxiety.
An important study in this field was conducted by Glasspool, Fox, Coulson, and Emery (2001). As part of the Risk Assessment in Genetics (RAGs) project, they developed a computer-based tool for assisting general practitioners and patients in making decisions about genetic risk of cancer. They constructed the system by adopting an argumentation logic approach that assesses genetic risk during a consultation and provides advice in the form of explanations. Glasspool et al. emphasised the use of argumentation as a way to be more in line with how people naturally think about probability and possibility.

Another important study was conducted by Green (2006). In the perspective of helping lay audiences to understand biomedical arguments, Green illustrated a discourse generator that creates an intentional-level representation of arguments.

It has been proved that inducing active processing – i.e. generation, re-organisation and self-explanation of presented material - improves understanding and recall of the information provided (Cosmides and Tooby 1996; Wadey and Frank 1997; Natter and Berry 2005). The use of argumentation in the delivery of health-related information facilitates this active processing by presenting a contextualisation of the claims at stake, i.e. in our case, the level of risk and the options available (Rubinelli and Schulz 2006; Schulz and Rubinelli 2006, 2008). Argumentation offers motivators to individuals’ internal reasons (factual information and intentional states) and can help them in the appraisal of the relative weights of the whole set of motivators as a way to arrive at a decision. Argumentation stimulates deeper processing and more critical thinking over a piece of information (Voss 1991; Kuhn 1993; Zeidler 1997).

In light of this, we designed OPERA according to a conceptual framework which is argumentative in nature, but more refined than RAGs in the personalisation and format of the information it delivers. RAGs produces final explanations as lists of reasons in support of a claim. OPERA works on the basis of a text-generation matrix that combines data from medical knowledge bases and data on the specific patient (Cawsey, Webber, and Jones 1997), and reorganises them into sentences which are fluent and at a higher level of granularity (Bickmore, Giorgino, Green, and Picard 2006). OPERA combines the personalisation of the information – as done by programmes, such as STOP (Smoking Termination through cOmputerised Personalisation, Reiter, Robertson, and Osman 2003) – with an argumentative framework.

3. The analytical framework

Before entering the discussion of the communication strategy adopted for OPERA, we present a summary explanation of how its engine works (Rubinelli, Bolchini, Paolini, and Schulz 2007). The risk level and the appropriate management options run by OPERA are categorised according to the UK National Institute of Health and Clinical Excellence (NICE) guidelines published in 2004, on the basis of (i) the number of relatives on the same side of the family with the same cancer or cancers that are known to run together; (ii) the ages of these relatives at diagnosis and (iii) the closeness of the family relationship with the person who is doing the assessment.

OPERA invites users to enter details of their personal and family history of breast and/or ovarian cancer. It also gathers information on age, sex, ethnicity and history of other cancers that are relatively rare. The user is led through the process by a series of questions about his/her demographic data and family history situation (e.g. “Have you ever been diagnosed with breast cancer?” and “In your family, how many relatives on the same side of the family have had breast and/or ovarian cancer?”). As the user answers the questions, the engine determines which questions should be asked next and finally computes the level of risk via a decision-tree. The inner working of the engine is determined by a number of rules that have the conjunctive form:
IF (A and B and C and …) IMPLIES $a$; where $a$ can be:

- a question number (determining if a question should be asked or not);
- the level of risk (i.e. low, medium or high risk) the user is grouped into;
- a piece of information (determining if that piece should be delivered to the user or not).

After the test ends, the feedback conveyed to users comprises two basic elements, which are joined to formulate the information package, namely:

- the summary: a textual expression of the most relevant data to assure the correctness of users’ answers;
- the risk assessment message.

Given all this, OPERA implements an explanation message that justifies the information it provides. The overall message it delivers is structured according to a specific model of argumentative reasoning, that is Toulmin’s (1958) theory of reasoning. As shown elsewhere (Green 2006), this theory allows an identification of all the elements that are needed both to justify a claim and to assess its strength. Central to Toulmin’s theory is a six-element structure that can be used to generate arguments, namely:

1. **Data**: The evidence, facts, background and information on which a claim is made.
2. **Warrant**: The knowledge that justifies a claim made using the data.
3. **Backing**: The knowledge material that validates the warrant.
4. **Qualifier**: The verbalisation of the confidence with which the claim is supported to be true.
5. **Rebuttal**: An exception of aspects for which the claim presented is not true.
6. **Claim**: The assertion or conclusion put forward for general acceptance.

Figure 1 shows the meta-argument structure of OPERA.

![Figure 1. A meta-argument structure for OPERA, based on Toulmin’s argument structure. The structure reads: Given Data, therefore Claim, since Warrant, because Backing, unless Rebuttal.](image-url)
The above scheme results in the information package delivered by OPERA composed of five main parts, namely:

1. **Your present situation** (the *Claim*, part a): an indication of the risk level.
2. **Explanation** (the *Backing* and the *Warrant*, on the basis of user *Data*): the justification of "Your present situation" on the basis of the data inserted and the NICE guidelines.
3. **What next?** (the *Claim*, part b): a description of possible options which may be offered and why.
4. **What might change your current situation?** (the *Rebuttal*): a general warning stating that, should users discover new information about their personal or familial history, they have to redo the assessment.
5. **How confident can you be?** (the *Qualifier*): a set of information on the "authority" of Cancer-backup and the NICE guidelines.

To quote an example of how this structure is implemented, we present in Figure 2 the message that OPERA delivers to a user with the following characteristics:

Currently aged between 19 and 39, female, not had breast cancer, not had ovarian cancer, no men in the family who have had breast cancer, one female relative diagnosed with breast cancer when she was over 40, no Jewish ancestry, no history of some unusual type(s) of cancer in the family.

Shifting to the specific contents of the argumentation, we applied a deductive model of argumentation "by association". The term argumentation by association goes back to the New Rhetoric (Perelman and Olbrecht-Tyteca 1958); it indicates those argumentative moves where two elements that are separate in the mind of an audience are coupled in light of a certain (more or less) explicit relation. Considering that for the New Rhetoric there are different types of argumentation by association, we chose as the main frame for OPERA, an argument by association on "the structure of the reality". By using as premises "facts" (propositions about reality that can be assumed without further justification, in the context of OPERA information about users’ personal and familial history) and "truths" (propositions that make connections about facts, in our case the NICE guidelines) we designed a message where the main claims (i.e. *You are at risk X* and *You should do Y*) are presented as the effects of the premises, with a causal link aimed at transferring the audience’s acceptance of the cause to the effect. Overall, the validity of this link is explicitly linked to its source of provenance – the NICE guidelines – that, as such, provides an argumentative input from authority to the factual contents delivered by OPERA.

The reasons behind the risk-level are, then, offered by means of a tailored explanation based on the selection and combination of those attributes that are particularly relevant for supporting the diagnosis of risk in light of the NICE guidelines. Thus, in our example, the explanation of the above risk is justified by pointing out the small number of relatives who have a history of breast cancer, as well as the age at which the one family member with cancer was diagnosed.

In a second stage, we decided to include a rhetorical (persuasion-oriented) component in OPERA’s argumentation that aims at assessing and addressing low-risk users’ eventual concerns. Indeed, one of the key premises from which argumentation theory proceeds is that influencing real audiences is not simply a matter of presenting facts and premises which are scientifically validated. To enhance persuasiveness, it is important to take into account values, attitudes and opinions that people might have (van Eemeren 2010). As discussed by Bennett et al. (2008), about 25% of people that undertake cancer genetic risk assessment based on family history show high level of distress. The main factors that are likely to influence an emotional response to a risk diagnosis include people’s understanding of the nature of the health threat and their expectations of its outcome. A deeper personalisation of the argumentation appears here as a necessary requirement.
1. YOUR PRESENT SITUATION
Your answers suggest that it is unlikely that there is a faulty breast cancer gene in your family. This is because you stated that you have only one female relative who had breast cancer and she was over 40 when the cancer was diagnosed. As you will see in the explanation below, the number of relatives with cancer as well as age at diagnosis are important factors when considering inherited risk of cancer.

2. EXPLANATION
The risk of breast and ovarian cancer being caused by an inherited faulty gene is assessed by considering your family history of breast and/or ovarian cancer. The following are all important considerations:

- the greater the number of relatives on the same side of the family with either of these cancers
- the more closely related they are
- the younger the age at diagnosis

Factors such as Jewish ancestry and some rare cancers developing at a young age are also taken into account. Research has so far identified two main breast and ovarian cancer genes, called BRCA1 and BRCA2 (Breast Cancer 1 and 2 genes). These genes have been linked with both breast and ovarian cancer. This is why the occurrence of either cancer in the family is important when considering inherited genetic risk.

Inherited changes in the BRCA genes (called genetic mutations) which would make a person more prone to developing breast or ovarian cancer, are usually only tested for in families where there is a very strong family history of those cancers. Exceptions may sometimes be made in families with Jewish ancestry or in families where there is a history of unusual cancers which were detailed in one of the questions you were asked, so these factors are particularly important to note if they apply to you.

It is important to remember that only a small proportion of cancers are caused by inheriting a faulty gene. The majority of cancers occur as a result of gene changes that happen by chance during a person’s lifetime.

3. WHAT NEXT?
If you are concerned about breast or ovarian cancer occurring in your family, you should discuss your situation with your doctor. In this section you will find information about breast screening, ovarian screening and genetic testing. However, based on your answers to the questions in this program, unless an exception is made, it is probably unlikely that your family will be offered genetic testing or extra screening on the NHS. Other risk factors can play a bigger role than family history in the development of breast or ovarian cancer and you can find information about those in this section too.

Breast screening
Unless your family history changes or an exception is made, it is unlikely that you or members of your family will be offered extra screening for breast cancer on the NHS. However, all women who are registered with a GP, are automatically invited to join the NHS Breast Screening Programme, once they reach the age of 50.

Women who do not have a strong family history of breast cancer are not routinely offered screening on the NHS before the age of 50.

4. WHAT MIGHT CHANGE YOUR PRESENT SITUATION
Your personal assessment is based on your response to the questions in this program. Therefore any change in your personal or family history that would result in a change to your answers might have an impact on this and you are advised to repeat the assessment.

5. HOW CONFIDENT CAN YOU BE?
It can never be guaranteed that you or other members of your family will not develop breast cancer. All women are at risk of this disease and inherited genetic risk is just one factor in the list of known risk factors. Risk increases with age. That is why it is important for all women to join the NHS Breast Screening Programme at 50 and think about lifestyle factors that may help to reduce risk. Your personal assessment as presented here is in relation to your personal and family history alone and does not take into account any lifestyle factors. Your personal assessment is based on the guidance on familial breast cancer that was issued to the NHS by the National Institute for Health and Clinical Excellence (NICE). NICE provides national guidance on the promotion of good health and the prevention and treatment of ill health.

Figure 2. Example of a risk assessment message delivered by OPERA to a low-risk user.

for understanding the type of concerns that users might have and meeting the range of their eventual expectations.

We considered the above assumption to be particularly relevant in dealing with the low-risk group. People in this category of risk do not have a significant personal or family history of cancer. Yet, they must have a reason for utilising the programme, including, as mentioned earlier, the possibility that they perceive themselves in a higher risk category. We decided to address this reason to prevent potential resistance to good news. The phenomenon of risk perception resistance is, indeed, known in the literature (Dillard, McCaul, Kelso, and Klein 2006).
To help the assessment of users’ motivations to access OPERA, we screened possible sources of influence on their risk-perception, considering what could influence the perceived threat of a hereditary cancer link. By moving from a social-interaction perspective, we assumed that individuals’ beliefs and opinions are the product of micro-individual characteristics and of a social construction activity performed within the social networks of the single person. Following the indications by the genetic counsellors who collaborated in this project, and have years of experience with the concerns that people express to Cancer-backup, we decided to consider three main possible categories of influence:

1. from healthcare workers;
2. from relatives and friends;
3. from the media.

In the outcome, we added a question at the end of the test for the low-risk group that enquires about the following sources of concern:

You decided to use Cancer-backup’s OPERA because:
• You recently read or heard something in the media about cancer in families.
• One or more relatives have had breast or ovarian cancer and you are worried about yourself and/or other family members developing it.
• You have had breast or ovarian cancer and you are worried about your children or other family members developing cancer.
• A friend has recently been diagnosed with breast or ovarian cancer.
• Someone in the health service suggested that you use this programme to find out about your risk.
• None of the above reasons.

The answer is then addressed in the tailored assessment. Going back to our example, if the user declared concerns due to family history, a section of the assessment argues against this concern in the following way:

It is nevertheless understandable that having a relative with cancer has caused you concern. The following information will hopefully help you to address your concerns and understand that your family history is not significant enough to indicate a higher risk.

The message continues by explaining how the risk is calculated and what aspects count into it:

The risk of breast and ovarian cancer being caused by an inherited faulty gene is assessed by considering your family history of breast and/or ovarian cancer. The following are all important considerations:

• The greater the number of relatives on the same side of the family with either of these cancers
• The more closely related they are
• The younger the age at diagnosis
• Factors such as Jewish ancestry and some rare cancers developing at a young age are also taken into account.

Through an essentially enthymematic (or elliptical) argumentation, the user is invited to make the inference that having only one relative who has had breast cancer (and, moreover, over 40) is not significant enough to indicate any possible genetic link.

Overall, the personalisation of the risk assessment is manoeuvred between an essentially deductive argumentation and, for the low-risk target group, a more dialogical component that has the rhetorical goal to enhance the credibility of the message.
4. Pilot-study

4.1. Study design

In this section, we present some indicators of users’ acceptance of OPERA. Data were collected through two self-administered questionnaires 1 month and 3 months after using OPERA during the period between March and July 2008. A non-stratified convenience sample of 61 enrollees aged ≥18, all women, was recruited through advertisements about OPERA on Cancer-backup’s website, two London free newspapers and a national newspaper. Eligible participants had to be able to read and speak English, to use a computer and have internet access. No other inclusion or exclusion criteria were used because the intervention was aimed at the general population. Enrolment took place between January and February 2008. Patients were asked to participate only after hearing a description of the project and providing written consent. They were asked to access OPERA in their own homes via a password. Drop-out rate between the first and the second questionnaire was 22% (N = 13). The study was approved by the London Medical Research Ethical Committee.

4.2. Results

On a general level, users were asked to indicate on a seven-point scale (1 = very much regret; 7 = very glad) whether they regretted using OPERA or were glad they had done that. Users were more or less all glad they had used OPERA, a month later. That is to say, almost all of them chose scale points on the “glad”-side of the scale. A majority chose even the most extreme point, indicating they were very glad. Some of this enthusiasm wears off after 3 months, but the gladness remains (Figure 3).

Another indicator of general evaluation is the importance ascribed to using the instrument. For many users, running through OPERA was an important thing to do (Figure 4).

Some effects of using the tool were measured by a question: “Here are some statements about the personal assessment. Please indicate on the scales underneath how much you agree or disagree with each statement”. Scales ran from 1 = I completely disagree to 7 = I completely agree. There was strong disagreement to a statement that one felt worried after reading the personal assessment. In contrast, respondents significantly agreed on that they felt less worried after using OPERA.

![Figure 3](image-url)  
*All in all, are you glad you used the risk assessment program or do you regret it?*

Question wording: “All in all, are you glad you used the risk assessment program or do you regret it?”.

Figure 3. Respondents are and remain glad they used OPERA.
Figure 4. Using OPERA was an important thing to do.

Figure 5. Assessment of OPERA after three months – worry and relevance.

OPERA and that others would also worry less after doing that (Figure 5). There is also strong agreement that OPERA is a good basis for making decisions to seek further advice on breast cancer (Figure 5).

Most respondents judged their family risk for breast cancer either as very low or very high. Respondents clearly tended to choose either one of the extreme scale points on this question, suggesting that the message sent by OPERA is rather clear and does not create confusion (Figure 6).

Forty-nine of the 61 users in this analysis could be classified according to the risk of being afflicted with hereditary breast cancer. Twenty-four respondents ran a low risk, five a moderate risk and 20 a high risk. Leaving aside, for sample size reasons, the moderate risk users, how did high-risk users and low-risk users react to the instrument?
Question wording: “Having used the program four or so weeks ago and anything that you have done since, how would you rate the risk of hereditary breast cancer occurring in your family?”

One month after test run, write-in interview, 61 respondents

Figure 6. Respondents’ assessment of hereditary breast cancer risk in family after having used OPERA.

Table 1. Recall of OPERA’s assessment of users’ risk.

<table>
<thead>
<tr>
<th>Risk of hereditary breast cancer</th>
<th>Low(%)</th>
<th>High(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>It is unlikely that there is a faulty breast/ovarian cancer gene in family</td>
<td>80</td>
<td>–</td>
</tr>
<tr>
<td>There is a slightly increased chance that there is a faulty breast/ovarian cancer gene in family</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>There may be a faulty breast/ovarian cancer gene in family</td>
<td>8</td>
<td>77</td>
</tr>
<tr>
<td>Do not recall</td>
<td>4</td>
<td>15</td>
</tr>
<tr>
<td>No answer</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>Total (n = 24)</td>
<td>100 (n = 20)</td>
<td></td>
</tr>
</tbody>
</table>

Note: Question wording: “Please tick which of the statements below was used in your personal assessment. If you do not recall reading any of the statements or would not prefer to answer this, please tick (d) or (e)”.  

As much as they remember what the personal assessment said, users recalled the gist of the message correctly. However, two users in the low-risk group indicated they were told there may be a faulty gene in the family (Table 1).

High-risk users mostly estimated their families’ risk of carrying the faulty gene as high or very high, and low-risk users as low or very low. That is to say: the messages sent by OPERA to the two risk groups reach their destinations. Generally, the assessment received is equally well understood and rated by high- and low-risk users. This conclusion holds true despite the fact that high-risk users tended to indicate in a clearly larger share that they did not learn anything from the personal assessment. The difference is not significant, though (Table 2). As most respondents who did not learn anything said they knew already, it can be assumed that knowledge of hereditary breast cancer and of their personal risk was not new to many of our high-risk users.

That high-risk users learned less is also evident from a question inquiring whether OPERA has changed a user’s level of understanding of hereditary breast cancer (Figure 7).

As we pointed out in the introduction of this article, OPERA should help users shape a realistic understanding of the risk. It should stimulate high-risk users to go for further tests, while reducing
Table 2. Learning from the personal assessment.

<table>
<thead>
<tr>
<th>Risk of hereditary breast cancer</th>
<th>Low (%)</th>
<th>High (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yes, I learned something from the personal assessment</td>
<td>71</td>
<td>47</td>
</tr>
<tr>
<td>No, have not learned anything</td>
<td>29</td>
<td>53</td>
</tr>
</tbody>
</table>

100 (n = 24) 100 (n = 19)

Notes: \(\chi^2 = 2.443, \text{df} = 1, p = 0.12\). Question wording: “The personal assessment you received in the end – do you think you learned something from that?”.

Figure 7. Change in level of understanding hereditary breast cancer.

Three months after using the tool, 11 in 17 low-risk users agreed (Points 1–3 on a 1–7 agree/disagree scale) to the statement “Having used the risk assessment program, I feel less worried about breast cancer”. Only three low-risk users disagree to that. Among high-risk users, most chose the middle point on the scale, two agreed and three disagreed.

The value of OPERA as an instrument for decision-making, claimed by the majority of users 3 months after the test, finds some specific highlights in the observation of some communication behaviours. A considerable share of high-risk users sought their doctor’s advice on hereditary breast cancer, or intended to do that, while low-risk users did not report having seen their doctor or intending to do so (Table 3 and Figure 8). This can indicate that low-risk users were satisfied with their personal assessment and not in need of further medical advice.
Table 3. Interpersonal communication behaviour after using OPERA.

<table>
<thead>
<tr>
<th>Risk of hereditary breast cancer</th>
<th>Low (%)</th>
<th>High (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Have (once or more often) talked to a doctor or a healthcare professional about hereditary breast cancer risk</td>
<td></td>
<td></td>
</tr>
<tr>
<td>One month after using the tool ($c^2 = 2.712, df = 2, p = 0.26$)</td>
<td>4 ($n = 24$)</td>
<td>16 ($n = 19$)</td>
</tr>
<tr>
<td>Three months after using the tool ($c^2 = 7.239, df = 2, p &lt; 0.05$)</td>
<td>$- (n = 17)$</td>
<td>36 ($n = 14$)</td>
</tr>
</tbody>
</table>

Note: Question wording: “Since using this assessment program, have you talked to a doctor/healthcare professional about hereditary breast cancer risk?”.

Figure 8. Likelihood of seeking doctor’s advice in future.

5. Discussion and conclusion

In this study, we have created a personalised risk-assessment message that presents an argumentative structure.

The promising evaluation by users in our pilot study points to some value in the communication model applied. At this stage, we can hypothesise the positive impacts on individuals’ risk-communication processing of two inter-related factors:

1. the addition of information content on risks;
2. the appeal to the reasoning faculty of users.

Previous studies have suggested that adding information to help people better understand the real extent of a risk positively impacts people’s cognitive outcomes (Hoopwood et al. 2001; Bowen, Burke, McTiernan, Yasui, and Andersen 2004; Edwards et al. 2008). With OPERA, we have focused on the impact of specific contents, i.e. the premises in support of the classification into a risk group and the available options. These premises provide a contextualisation of the risk in a way that can invite users to reflect on the objective (medically based) reasons behind it.
The appeal to reasoning is the second main factor on which OPERA rests. Considering the role of NICEs’ guidelines in the orchestration of OPERA’s explanation, we can say that the tool instantiates a deductive argumentation that is, ultimately, based on authority. Indeed, we expect users to recognise two main premises.

The focal premise is that the source of the message (i.e. Cancer-backup) is a genuine expert in cancer information. The second premise is that this source asserts propositions that address the clinical genetic history and the eventual concerns of the person being assessed.

The application of this argument scheme in the design of OPERA has been facilitated by a series of factors. Generally speaking, an argumentative structure like OPERA can be instantiated only if it can answer to the critical questions that are appropriate when evaluating an argument from authority (van Eemeren and Grootendorst 2004).

First of all, the institution acting as a source has to be recognised as an expert in the field. Given the national and international recognition of Cancer-backup, this aspect was less problematic.

Second, problems might arise when there is disagreement among several experts over an issue. When disagreement occurs, it is difficult for a layperson to arrive at a reasoned opinion or decision about the issue. Here, again, being OPERA based on the NICE guidelines, the coherence of the body of knowledge offered is guaranteed. Doubts might arise on whether the programme, being online, is used by non-UK people from countries that apply different national guidelines.

Another important factor is that the interpretation of the expert’s opinion by the layperson has to be carefully guided. To be useful, an expert opinion has to be in a form that is clear and intelligible. But, especially in dealing with genomics, the jargon can be technical and resists translation into accessible layman’s terms. We had to face the issue of how to transmit clinical guidelines to the lay people. With OPERA, we answered to this aspect by designing the wording together with genetic counsellors who had a live experience on the way people react to their words and their letters during or after consultation.

Further studies will have to address some of the important limitations of this study. First of all, in the recruitment of our sample, we did not fully take into account users’ previous knowledge. In our analysis we relied on users’ declaration on whether or not they have learned from the test. More effort is needed to specify the impact of OPERA according to different levels of pre-existing knowledge on hereditary breast cancer and personal risk.

Another limitation is that in the study, we did not evaluate the contribution of argumentation to the desired outcome, an aspect that, however, we plan to address through more refined experimental design and after the revision of the pilot version of OPERA.

Finally, the argument model used in this current version of OPERA is that all knowledge in the program is stored as pre-compiled arguments, assessed with an a priori screening of users’ informational needs. This is an approach that has the clear advantage of being simple, but it presupposes an anticipation of all possible arguments.

At this stage, we have worked on the assumption that the authority of the scientific guidelines is sufficient enough to meet the range of users’ emotional concerns. More complex argumentative structures, performing a higher level of dialoguing interaction with users, are difficult to implement.

We are currently exploring how theories of argumentation based on the critical discussion of arguers – in particular, the pragma-dialectical theory developed by van Eemeren and Grootendorst (2004) and current models of persuasion (O’Keefe 2002) – can advance theory and practice of this phenomenon. One promising attempt has been made by Grasso, Cawsey, and Jones (2009) who have formalised an agent, Daphne, that advises on controversial issues by using dialectical argumentative strategies. Daphne, however, does not recognise the opponent’s argumentative schema and this risks limiting the actual resolution of eventual differences of opinion.
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References


