

Table 2 Multivariate logistic analysis of risk factors for preterm delivery in Denmark, 1995-2004

	All preterm		Extremely preterm*		Very preterm*		Moderately preterm*		
	No	Odds ratio (95% CI)	No	Odds ratio (95% CI)	No	Odds ratio (95% CI)	No	Odds ratio (95% CI)	
Mode of conception:									
Spontaneous	63 0928	34 039	1.00	1379	1.00	3607	1.00	29 053	1.00
In vitro fertilisation	15 216	2 711	1.25 (1.19 to 1.32)	197	1.76 (1.47 to 2.12)	345	1.15 (1.01 to 1.31)	2 169	1.17 (1.10 to 1.24)
Initiation of delivery:									
Spontaneous	55 7267	29 870	1.00	1433	1.00	3344	1.00	25 093	1.00
Elective	88 877	6 880	1.09 (1.06 to 1.12)	143	0.39 (0.33 to 0.46)	608	0.77 (0.71 to 0.85)	6 129	1.19 (1.15 to 1.23)
Parity:									
Primiparous	28 7080	19 938	1.53 (1.49 to 1.57)	931	1.79 (1.60 to 1.99)	2240	1.63 (1.52 to 1.74)	16 767	1.48 (1.44 to 1.51)
Multiparous	35 9064	16 812	1.00	645	1.00	1712	1.00	14 455	1.00
Type of pregnancy:									
Singleton	633 012	30 859	1.00	1250	1.00	3171	1.00	26 438	1.00
Multiple	13 132	5 891	14.03 (13.75 to 14.90)	326	11.12 (9.6 to 12.87)	781	11.88 (10.83 to 13.02)	4 784	11.70 (11.22 to 12.20)
Maternal age†	646 144	36 750	1.01 (1.01 to 1.01)	1576	1.03 (1.02 to 1.04)	3952	1.02 (1.01 to 1.02)	31 222	1.01 (1.01 to 1.01)
Year‡	646 144	36 750	1.03 (1.02 to 1.03)	1576	1.06 (1.04 to 1.07)	3952	1.03 (1.01 to 1.04)	31 222	1.02 (1.02 to 1.03)
Smoking:									
Non-smoker	489 129	24 820	1.00	916	1.00	2479	1.00	21 425	1.00
Smoker	131 914	9 190	1.52 (1.49 to 1.55)	379	2.26 (2.11 to 2.42)	1041	1.76 (1.68 to 1.85)	7 770	1.41 (1.38 to 1.44)
Ethnic origin:									
White European	608 339	34 778	0.93 (0.88 to 0.97)	1493	0.86 (0.69 to 1.07)	3751	0.94 (0.81 to 1.09)	29 534	0.94 (0.89 to 0.98)
Other	37 797	1 972	1.00	83	1.00	201	1.00	1 688	1.00

CI, confidence interval.

All odds ratios were mutually adjusted for other variables in the table.

*Extremely preterm: 22-27 completed weeks of gestation; very preterm: 28-31; moderately preterm: 32-36; term: ≥ 37 .

†Odds ratio for a one year change in age.

‡Odds ratio for a one year change in calendar time.

different proportions of high risk pregnancies and parity, but using standard populations of primiparous women at low risk identified by the criteria described here could enable valid international comparisons of spontaneous preterm delivery rates to be made.

Contributors: JL-R conceived the study in collaboration with UK. SR retrieved register data and performed the initial analyses. UK performed additional statistical analyses. All authors contributed to the data interpretation. JL-R wrote the first draft of the manuscript and all authors contributed to the revision.

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Accuracy and self correction of information received from an internet breast cancer list: content analysis

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Abstract

Objectives To determine the prevalence of false or misleading statements in messages posted by internet cancer support groups and whether these statements were identified as false or misleading and corrected by other participants in subsequent postings.

Design Analysis of content of postings.

Setting Internet cancer support group Breast Cancer Mailing List.

Main outcome measures Number of false or misleading statements posted from 1 January to 23 April 2005 and whether these were identified and corrected by participants in subsequent postings.

Results 10 of 4600 postings (0.22%) were found to be false or misleading. Of these, seven were identified as false or misleading by other participants and corrected within an average of four hours and 33 minutes (maximum, nine hours and nine minutes).

Conclusions Most posted information on breast cancer was accurate. Most false or misleading statements were rapidly corrected by participants in subsequent postings.



Details of false or misleading statements are on bmj.com

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Introduction

Nearly half of women recently diagnosed as having breast cancer turned to the internet for information on health.¹ Consumers are satisfied with their online experience and are making choices based on the information that they encounter.²⁻³ In some cases patients may not discuss with clinicians their use of treatments found online.⁴ On the other hand, clinicians are faced with patients who have been informed (or misinformed) by information posted on the internet. Clinicians, researchers, and healthcare consumers are therefore concerned about the accuracy of online health information.²⁻⁵⁻⁷

Internet cancer support groups offer the public a forum in which to share experiences, ask questions, and offer advice. Discussions have been categorised according to their content as related to quality of care, treatment, recurrence of disease, and alternative therapy.⁸ Rates of false or misleading information may vary depending on the definition of falsehood (for example, verified by multiple observers, possibly or definitely false?), forum studied, existence of a moderator, prevalence of health professionals, topic, and other factors. One study found that about 6% of postings to an online epilepsy forum were “objectively inaccurate.”⁹ A review of a German language brain tumour list found that “Normally, precisely formed questions were answered by medically correct and solid statements. When medical statements were incorrect, other participants did not hesitate to correct or criticise.”¹⁰

Professionals rely on the peer review process to screen out false or misleading information. Peer reviewed journals ensure that published articles are critically reviewed by experts. Similarly, open source software relies on the user community for quality control. A comparable peer review process may occur on online forums whereby false or misleading information is corrected quickly and reliably by subsequent postings. This possibility, which may be referred to as the “self correction hypothesis,” has been suggested in the literature.⁹⁻¹¹⁻¹² To our knowledge the self correction hypothesis has never been quantitatively tested in an unmoderated forum. We determined the prevalence of false or misleading statements found on a specific internet cancer support group site and whether these statements were identified and corrected by subsequent postings.

Methods

We selected the Breast Cancer Mailing List because it is unmoderated and therefore reflects only the opinion of voluntary participants without any kind of systematic interference from healthcare professionals. The list was formed in 1994 and facilitates communication between people affected by breast cancer.

Participants include individuals with breast cancer; their caretakers, family, and friends; and a small number of healthcare professionals. At the time of our analysis, there were about 500 participants to the list, with an average of 310 distinct people uploading messages per month.

A person can join the Breast Cancer Mailing List by sending an email message to the mailing list administrators. After receiving a message of confirmation, the participant then receives all messages (postings) sent to the list’s email address. The participant can post to the list using the same address. The list’s archives contain all messages (postings) since its inception. At the time of our study the archives contained over 600 megabytes of information in 248 051 text email messages and were publicly available at <http://bclist.petebevin.com/>

We reviewed all messages posted to the list from 1 January to 23 April 2005. A “post” or “posting” was defined as the entire content of a message sent by a participant to the list. We reviewed the entire posting to identify false or misleading statements. A “thread” was defined as a group of postings under one subject. A thread started with the first posting of a new subject and ended when no more postings occurred in response to that subject. The postings in a thread were in chronological order. We excluded from our study postings that were incomplete at the cut-off date (23 April).

Three independent clinicians reviewed and classified the postings: a general practitioner (AE), a general internist (EVB), and a breast cancer surgeon (FMB). The general practitioner first identified statements that might be false or misleading. Factually incorrect statements and those that were likely to lead a medically naive reader to a false factual conclusion were defined as false and misleading, respectively. For the purposes of this study we did not consider any statement that was phrased as an opinion (I believe that . . .) or a question to be false or misleading. We did not require an explanation of why the original statement was incorrect to consider the false or misleading statement successfully identified and corrected. However, the correction had to relate to the specific statement that was false or misleading and give the correct facts. We recorded the location of the posting within the thread, the date and time the message was posted, and the number of postings until a participant identified the incorrect information. We considered a false or misleading statement as not identified when we found no evidence of its identification by other participants and the thread was exhausted.

EVB and FMB verified the information in the candidate postings identified by AE and validated or challenged the initial judgment. Reviewers EVB and FMB confirmed or dismissed all possibly false or misleading postings, based on review of relevant literature. Differences were resolved by consensus.

Results

Overall, 4600 postings, organised into 1378 threads, were reviewed. Table 1 shows a summary of the total number of postings reviewed per month. AE identified a total of 32 (0.7%) candidate statements containing false or misleading information in 30 threads. Of these

Table 1 Number of threads and postings reviewed

Variable	January	February	March	April*	Total
No of threads	448	328	359	243	1378
No of postings	1487	1131	1197	785	4600
Average No of postings per thread	3.32	3.45	3.33	3.23	3.34
No of authors	308	313	308	313	

*Cut-off point of study was 23 April.

What is already known on this topic

Healthcare consumers search the internet for information on health

Online information affects patient's decisions about treatment

Despite the publication of many quality measures, no validated, usable measures exist that can reliably identify false or misleading information online

What this study adds

Given a sufficiently active forum, participants can identify and correct most false or misleading statements quickly and reliably without requiring professional review

Online forums can police themselves

32, 10 (31%) were confirmed to be false or misleading by the other two reviewers on the basis of the relevant literature and consensus among the three independent reviewers. The statements were posted by seven different participants, none of which identified themselves as health professionals. A list of the statements confirmed as false or misleading is on bmj.com.

Table 2 summarises the participants' level of activity before and during the study period. The participants who had posted or corrected false or misleading information were among the historically most active users of the list. On average these nine participants ranked among the top 1.6% of participants based on the number of postings and in the top 9.5% of participants during the study period.

Seven of the 10 false or misleading statements were identified and corrected by six participants, three of whom had also posted false or misleading statements. The average time before a false or misleading statement was identified was 4 hours and 33 minutes from when the posting appeared on the forum (range 2 h 3 min-9 h 9 min). For the false or misleading statements that were identified and corrected, the average number of postings before identification was 2 (range 1-3).

Discussion

Few postings on the internet cancer support group Breast Cancer Mailing List contained false or misleading information. If false or misleading information was posted to the mailing list, it was identified and corrected by a subsequent posting in more than two thirds of cases. Multiple participants posted and corrected false statements, rather than a single expert participant. One individual did, however, post three of the seven corrections that related to a single topic (expiry of drugs).

In contrast to our results, one study found that conclusions drawn by authors on a mailing list for painful hand and arm conditions were rarely questioned.¹³ This finding was not, however, quantified. One possible explanation for this discrepancy is that our study was carried out 11 years later. In 1994, the internet was

Table 2 Participants' level of activity before and during study period

Participant	No of false or misleading postings		Historical level of activity (12 430 participants)		Level of activity during study (313 participants)	
	Produced by participant	Identified by participant	No of postings	Activity ranking	No of postings	Activity ranking
A	2	1	1051	16	51	29
B	2	1	772	30	64	23
C	0	1	272	148	175	4
D	1	0	190	207	18	54
E	2	1	188	208	186	2
F	1	0	128	261	61	25
G	1	0	85	303	48	31
H	0	1*	18	370	8	64
I	1	0	5	383	5	67
Mean			301	214	68.4	33.2

*This single posting corrected three related but distinct false or misleading statements, therefore total is 10 false or misleading statements, seven corrections.

relatively new and our findings may reflect a maturing medium in which participants are more likely to critically evaluate information. In addition, because participants may have already experienced phases of the disease, they can provide accurate information.¹⁴ Perhaps there is more shared experience among patients with breast cancer than among people affected by limb pain. In addition, most of the false or misleading statements that we identified were not likely to lead to harm.

Our study was limited by the fact that a single reviewer determined the statements that might be false or misleading. We may therefore have missed some false or misleading statements. For this reason we did not require that a statement be factually incorrect and allowed misleading statements, as defined above. Furthermore, reviewers were not blinded to the study hypotheses. An additional limitation of our study is that we analysed a single, albeit large, internet cancer support group. Our findings may not generalise to other online forums. Unlike most previous studies, however, we chose a large, unmoderated list that truly reflects self correction, rather than the knowledge of a moderator. More research is needed to determine if our findings generalise beyond the Breast Cancer Mailing List to other online communities and other health topics.

Ideally consumers would have access to accurate online information without direct professional guidance, so that the limited time they have with clinicians could be used more efficiently. This requires that online resources present accurate information. At this time, no known effective strategies exist to ensure that online information is accurate. Our findings suggest that, given a forum, the internet can police itself.

Contributors: AE, FM-B, and EVB planned the study, collected the data, and drafted and revised the manuscript. EVB is guarantor.

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Effect of different forms of information produced for cancer patients on their use of the information, social support, and anxiety: randomised trial

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Abstract

Objective To explore the hypothesis that different methods of selecting and printing information for cancer patients could improve emotional support by affecting interaction with others, and so lead to improved psychological wellbeing.

Design Randomised trial with eight groups (three factors, 2×2×2). Data collected at recruitment and three month follow-up.

Participants 400 patients starting radiotherapy, of whom 325 with breast or prostate cancer and complete anxiety and depression data were included in the analysis.

Interventions Printed booklets: half had only general information from CancerBACUP about each patient's cancer and half had personalised information from the patient's medical record plus selected general information; half were composed of information chosen interactively by the patient and half were produced automatically with a larger volume of material; and half had additional advice on anxiety management.

Main outcome measures Patients' views of the information, use of their booklets with others; change in reported social support; change in anxiety and depression.

Results The larger booklets produced automatically were more likely to be found useful and to tell patients something new and less likely to be seen as too limited than the booklets produced interactively, but they were also more likely to overwhelm some patients. Personalised booklets were more likely than general booklets to tell patients something new. There was no difference in patients' perceived understanding of their cancer by any of the intervention factors. Patients with personalised information were more likely to show their booklets to others and to think it helped in discussing their cancer or its treatment. There were no major

differences in social support, anxiety, or depression by any intervention factors.


Conclusions Patients were more likely to show personalised information to their confidants than general information. Further research is needed into the effects of sharing information on patients' social support and anxiety.


Trial registration US Government Clinical Trials Database NCT00127465

Introduction

Anxiety and depression are important and common comorbidities in cancer and may affect survival. Providing information and social support may improve patients' psychological wellbeing,^{1 2} but different patients may have different information preferences³ and coping styles.⁴ Patients fare better when the information they receive is tailored to their coping style. Coping style also affects patients' desire for social support, and availability of a confidant and a means by which a patient can engage support are also important.⁵

In a previous study we found that patients preferred personalised information to general information.⁶ Unexpectedly, we found that patients with personalised information showed better improvement in anxiety over three months than those with more general information. We knew that patients with personalised information were more likely to show it to someone at home and hypothesised that this might be partly responsible. We have now carried out a randomised trial, with similar patients and setting, to explore the hypothesis that different methods of selecting and printing information for cancer patients

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 Appendices 1-10, providing further information about the study, are on *bmj.com*