Uptake of Genetic Testing and Pre-Test Levels of Mental Distress in Norwegian Families with Known BRCA1 Mutations

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ABSTRACT: 232 family members from 27 Norwegian families with BRCA1 mutations were offered genetic testing. 180/232 (78%) chose to be tested, 14/232 (6%) have not yet decided and 38/232 (16%) declined. All 232 persons were invited to fill in the following questionnaires when offered testing: Impact of Event Scale (IES), Hospital Anxiety and Depression Scale (HADS), General Health Questionnaire (GHQ-28) and Beck Hopelessness Scale (BHS). 207/232 (89%) responded to the questionnaires. Of those declining to be tested 23/38 (61%) answered the questionnaires compared to 170/180 (94%) of those wanting the test (p < 0.0001). A higher proportion of females with a history of cancer than females without such a history had abnormal scores on the IES-intrusion and GHQ questionnaires (p < 0.001). Healthy females who were deciding on predictive testing had the same or lower prevalence of mental distress compared to the general population, between 4.3% and 18.0% as measured by the different questionnaires. Males did not differ from healthy females on any of the measures. According to their HADS scores, women without a history of cancer deciding on predictive testing for breast-ovarian cancer had lower or equal levels of mental distress compared to the general population. The high uptake of genetic testing combined with the lower than normal prevalence of mental distress indicates that the activity may continue as practised, awaiting longitudinal data concerning the levels of mental distress after genetic testing.

INTRODUCTION

The finding of founder mutations of BRCA1 in Norwegian families with heritable breast and ovarian cancer has made it possible to identify a number of mutation carriers, and offer genetic testing to their relatives. Previously these relatives were given genetic counselling, on the basis of clinical criteria, that gives female members in the same kindred an increased risk of contracting cancer. Genetic testing provides more certain knowledge, with both the possibility of good news (negative test) — risk comparable to that of the rest of the population — and the possibility of bad news (positive test) that increases the likelihood of contracting cancer. The advent of genetic testing makes it of interest to know more about recruitment to testing and how family members cope with this opportunity.

Previous reports have shown that interest in testing has been high but that a limited number of relatives actually go on to be tested [11], and a high number of family members report that they anticipate negative psychological reactions to a positive test [14]. In contrast to the public debate on assumed adverse effects, no significant psychological disturbances among the tested family members have actually been reported. A summary of the literature has been given [2]. Our knowledge of predictive genetic testing for adult-onset diseases is based largely on experience with incurable diseases, like Huntington’s disease.
review concludes that both carriers and non-carriers experience short-term emotional reactions, but no long-term reactions have been revealed [13]. Whether these findings are relevant for a different disease with lower penetrance and a possibility of cure, such as inherited breast cancer, has not been examined. As the first report in a longitudinal study concerning the psychological effects of genetic testing, we here report on the uptake of testing in Norwegian families with demonstrated mutations in BRCA1 who have been offered testing, their compliance with psychosocial questionnaires, their prevalence of mental distress and levels of anxiety and depression at the time when they are offered a test.

MATERIAL AND METHODS

The population studied is drawn from our cancer clinic cases, including more than 1000 breast cancer kindreds as clinically defined. Most of the families are self-referred, and we have demonstrated the underlying BRCA1 mutation in a number of families. We have earlier published our findings of founder mutations and the results of our program for early detection of breast and ovarian cancer [3,4,12]. As of December 31st 1998, 232 members of 27 families with demonstrated BRCA1 mutations had been offered genetic testing. 156 (67%) were women who had not had cancer, 46 (20%) were men (none had experienced cancer) and 30 (13%) were women who had had breast and/or ovarian cancer. As agreed at earlier genetic counselling sessions, they were offered genetic testing when it became available. At the same time as they were offered testing they received written information about the different aspects of testing; they were invited to pre-test counselling, and they were also invited to complete the psychosocial questionnaire described below before coming to counselling. One written reminder was sent to those not responding to the first request after 8 weeks.

Levels of anxiety and depression were measured with the Hospital Anxiety and Depression Scale (HADS) [7]. The questionnaire has 14 items, half of which reflect anxiety and half depression. HADS gives overall scores for anxiety (min 0, max 21) and depression (min 0, max 21). A score of 8 or higher on each sub-scale indicates possible “caseness”, and a score of 11 or higher definite “caseness”. We applied the lower threshold, to increase the sensitivity for possible “cases”.

The General Health Questionnaire (GHQ-28) is focused on psychosocial distress and subjective wellbeing. We used the “scaled version” which has 28 items divided into four sub-scales, social functioning, somatic symptoms, anxiety/sleep disturbances and depression. To identify cases we applied “simple” scoring, in which each item is given a score based on the symptom both being present and being present to a greater extent than usual (0, 0, 1, 1). The threshold is usually between 5 and 6, but it has been suggested to use a threshold between 7 and 8 when screening a population with a somatic disease [6,9]. We applied both in this study, since one subgroup had a history of cancer and the other two had not.

Degree of hopelessness was measured with the Beck Hopelessness Scale (BHS), a questionnaire that has 20 items. BHS scores can vary from 0 to 20, with scores of 9–13 indicating moderate hopelessness and scores ≥ 14 indicating severe hopelessness [1]. We applied the lower threshold of between 8 and 9 for “caseness”, in order to identify all possible “cases” with high sensitivity.

The Impact of Event Scale (IES) is a questionnaire that measures psychological distress on two sub-scales, “intrusion” that refers to intrusively experienced images, thoughts, feelings and dreams and “avoidance” that refers to consciously recognised avoidance of certain feelings, ideas or situations. The intrusion sub-scale has 7 items, the avoidance sub-scale has 8 items, and each item has a score of 0–5. A score on each sub-scale of 9–19 denotes a moderate level of distress and a score of ≥ 20 denotes more severe distress [8]. We applied the threshold of a score of > 20, because low to moderate stress levels were expected to be frequent in individuals offered genetic testing for an inherited disease.

All returned questionnaires with less than 90%
of each subscale properly answered, were excluded from analysis of that subscale. When less than 10% of the items were missing, missing items were given a score according to the mean of the answered items.

RESULTS

As of May 1st 1999, 180 out of 232 family members have received the results of genetic testing (78%), 14/232 (6%) have not decided yet and 38/232 (16%) have declined. Of 158 females without a history of breast and/or ovarian cancer 120 had opted for testing (76%).

Total compliance in answering the questionnaires was 207/232 (89%). Compliance among females without a history of cancer was 170/180 (94 %), than of those not wanting the test, 23/38 (61%) (p < 0.0001). This was also true for females without a history of cancer. 114/120 (95%) of those who wanted testing answered the questionnaires compared to 19/29 (66%) of those who did not want testing (p < 0.0001)

Details of distribution of scoring on each questionnaire are given in Table 1. For females without a history of cancer we found that clinical levels of mental distress varied from 6/142 (4.3%) on the HADS-Depression sub-scale to 26/142 (18.0%) on the HADS-Anxiety sub-scale. All results from the other questionnaires were within this range. For females with a history of cancer the proportion with mental distress varied from 3/25 (12.5%) on the HADS-Depression to 10/25 (41.7%) on the IES-Intrusion (IES-I) sub-scale.

There was a statistically significant (p < 0.01) higher number of cases among the females with a history of cancer compared with females without such a history, as measured by the IES-I and the GHQ-28. The number of cases was 10/25 (41.7%) versus 15/142 (10.7%) on the IES-I and 9/25 (37.5%) versus 18/142 (12.7%) or 11/142 (7.8%) on the GHQ, depending on the threshold value. There were no significant differences between males and females without a history of cancer on any of the questionnaires.

DISCUSSION

The uptake of genetic testing was as high as 78% in our study: Previous reports give uptake rates between 22% [5] and 43% [11]. A high uptake rate can be understood as a high degree of trust that inherited breast cancer may be cured, in confidence concerning the test results, and in the fact that Norway has a public health care system basically without cost for the patients. They are offered life saving health services for free, and their access to insurance/bank loans has not so far been affected. The high compliance rate in terms of return of completed

| Scale   | Cut-off | Females without cancer (n = 142) | Females with cancer (n = 25) | Males (n = 40) |
|---------|---------|----------------------------------|-----------------------------|----------------|
|         | Mean (SD) | "cases" n (%) | Mean (SD) | "cases" n (%) | Mean (SD) | "cases" n (%) |
| IES-I   | >= 20   | 8.7 (7.2) 15 (10.7) 2 | 17.4 (8.6) 10 (41.7) 1 | 6.8 (8.1) 3 (7.9) 2 |
| IES-A   | >= 20   | 8.2 (7.2) 14 (10.1) 3 | 11.9 (8.4) 5 (20.8) 1 | 8.1 (7.7) 3 (7.9) 2 |
| HADS-A  | >= 8    | 4.5 (3.8) 26 (18.0) 1 | 5.8 (3.6) 7 (29.2) 1 | 3.7 (3.5) 6 (15.4) 1 |
| HADS-D  | >= 8    | 1.8 (2.2) 6 (4.3) 1 | 3.0 (2.8) 3 (12.5) 1 | 2.4 (2.9) 4 (10.3) 1 |
| GHQ-28  | >= 6    | 2.3 (3.8) 18 (12.7) 0 | 4.7 (5.2) 9 (37.5) 1 | 1.9 (4.6) 5 (12.8) 1 |
| GHQ-28  | >= 8    | 2.3 (3.8) 11 (7.8) 0 | 4.7 (5.2) 9 (37.5) 1 | 1.9 (4.6) 4 (10.3) 1 |
| BHS     | >= 9    | 4.0 (2.6) 7 (5.1) 5 | 5.6 (3.5) 4 (16.7) 1 | 4.4 (2.5) 4 (10.0) 0 |

1p<0.01 compared with females without cancer
2e.c.: excluded cases due to incomplete answers
questionnaires in our study (89%) give us results that are more representative, compared to the relatively low compliance in other studies [2]. Our sample size makes variations due to small numbers unlikely. The results obtained are, therefore, considered representative for the population studied.

In a population study of 33,168 females and 29,328 males aged 20–89 years in Norway [Stordal et al. Submitted] HADS-Anxiety (≥ 8) was 17.3% among females and 12.1% among males. Mean score in females was 4.47 (SD 3.45) and mean score in males was 3.87 (SD 3.08). The prevalence of depression in the same study was 10.4% among females on HADS-Depression (≥ 8) and 10.8% among males. Mean score in females was 3.33 (SD 3.04) and mean score in males was 3.54 (SD 3.03). Comparing these results with the present study, we found a lower level of mental distress measured by HADS-Depression ($x^2 = 5.2$, $p < 0.025$), and no difference as measured by HADS-Anxiety ($x^2 = 0.1$, $p > 0.1$) among women without a history of cancer.

The high levels of “caseness” on GHQ-28 among females with a history of cancer could be caused by somatic symptoms, and by the fact that they are experiencing distress both from the history of cancer, the possibility of having given their children the mutation. The interpretation of mental distress caused by the cancer itself is in keeping with a previous report on breast cancer patients [15]. In a study of post-traumatic distress in primary breast cancer in Norway Tjemsland et al. [15] found medium scores on IES (9–19) in 37% (IES-I) and 46% (IES-A) and high scores (≥ 20) in 44% and 29% respectively at the time after diagnosis but before surgery. The results one year after surgery were medium scores (9–19) in 37% (IES-I) and 29% (IES-A), and high scores (≥ 20) in 9% and 10% respectively.

In summary, our findings show an apparent lack of adverse psychological reactions to our offer of predictive genetic testing in both males and females without a history of cancer, but raised levels of mental distress and number of cases among females with a history of cancer. We also found an uptake of testing much higher than in any previous report and this may be an effect of self-referral to our health-service. Whatever the explanation of these results, they are as hoped for, when considering the future of offering predictive genetic testing for curable forms of cancer to healthy members in families with demonstrated mutations. The questionnaires used have previously been validated both...
nationally and internationally, and the number
of responders was large enough to escape
uncertainty relating to variation in small
numbers. We conclude that the high uptake of
genetic testing and the lack of adverse mental
distress found on pre-test questionnaires,
answered with high compliance by healthy
family members indicate that the activity of
predictive testing may continue without undue
fear of adverse psychological effects in those
offered testing. Continuous follow-up of the
patients described will give results on both short
and long term effects of our intervention both
for those who choose to be tested and for those
who choose not to undergo predictive testing.

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