Genetics of early cancer detection behaviours in Australian female twins

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Early detection of cervical and breast cancers is an important component of women's health strategy. Screening programmes, health professional interventions and preventive behaviours such as breast self-examination provide the means to this end. Our twin study sought to identify the relative influence of environmental and genetic factors on liability to early cancer detection behaviours, including use of cervical smear tests, mammograms, and breast examination. Additive genetic and random environmental effects models gave the best, most parsimonious fit to the data for each early cancer detection behaviour. The heritability of liability to Pap smear use was 66%, mammogram use 50%, breast examination by a doctor or nurse 38% and breast self-examination 37%. Genetic influences were behaviour-specific; there was no evidence for a common genetic influence on the four behaviours. Potential covariates investigated included age, amount of contact between co-twins, educational level and personality traits such as harm avoidance, novelty seeking, reward dependence, neuroticism, anxiety, depression, self-esteem, perceived control, interpersonal dependency and ways of coping. None were significant. The study was carried out before the implementation of national screening programmes with media campaigns to increase participation rates. Hence follow-up investigation, including data on regularity of behaviours, would be informative

Keywords: health behaviour, cancer screening, genetic influences, twins

Introduction

Breast cancer is the most common cause of cancer death for Australian women, with a lifetime risk of 1 in 15. Screening allows for early detection of malignancy with the aim of reducing mortality rates, as most identifiable risk factors for breast cancer are not preventable. Evidence suggests that mammography is contributing to the reduction in mortality rates from breast cancer. Early detection of breast cancers allows for considerable treatment cost saving compared with later detection. Cancer of the cervix is a less common but still significant health problem for Australian women. Early detection and treatment of pre-malignant changes to the cervix can prevent cancer of the cervix. This prevention strategy is based on cervical (Papanicolaou or Pap) smear examinations which can detect pre-malignant cell changes. Pap smear testing has been available on an opportunistic basis in Australia since the 1960s, following in the wake of its introduction in countries such as Britain.

We describe a study instigated in 1988, when there were no commonly accepted guidelines on cervical screening frequency in Australia. The Australian National Health and Medical Research Council had recommended three-yearly screening from the start of sexual activity for life (NHMRC 1984), whilst other bodies such as the American College of Obstetrics and Gynecology and the International Academy of Cytology recommended annual screening from age 18 or start of sexual activity for life. The first Australian guidelines were agreed on in July 1988. The National Program for the Early Detection of Breast Cancer and the National Policy on Screening for the Prevention of Cervical Cancer were initiated in 1990. In 1991 Australia introduced the Organised Approach to Preventing Cancer of the Cervix to implement the national policy. Because of Australia’s division of responsibilities between Federal and State Governments, determined by the Constitution, substantial variation exists between States in their level of implementation of national policies. Variability prevailed also prior to the national policy introduction. Nurse practitioners had been performing Pap smears and breast examinations since 1976 in the State of New South Wales, whereas in the State of Queensland these services were only available from medical practitioners.

The 19% fall in the number of women who died from cervical cancer in Australia between 1973 and 1993 has been attributed largely to Pap smear
had never had one.10 Concern has been consistently had a smear test for eight years or more, and 10% had never had one.10 Concern has been consistently expressed that women identified as being at high risk of cervical cancer are not utilising available screening services. Surveys of women who have presented to physicians with invasive cervical cancer have shown that over 60% of the women had never undergone a cervical smear test, and this proportion was highest in the older women.11

In 1989–90, 71% of women aged 18–64 years had had a Pap smear in the previous three years, with the highest rate (86%) among women aged 25–34 years.1 Women over 40 with the highest risk of cervical cancer had a lower screening rate than younger women at lower risk.5 Although cervical screening has contributed to reduction in incidence and mortality due to cervical cancer in many countries,12 there is some question about its role in relation to certain groups. Reports have suggested an increasing incidence of early stage cervical intra-epithelial neoplasia (CIN) in younger women,13 possibly due to the prevalence of human papilloma virus. Mortality rates have been rising amongst young women in Australia.12

Public health campaigns promoting use of screening procedures are costly,14 underlining the importance of understanding what lies behind use of early cancer detection behaviours (ECDBs). Cancer prevention directed behaviours may include general activities such as dietary modification. Other behaviours are specific in intent and purpose. These may be private activities; for example, women are encouraged to practise monthly breast self-examination to detect breast changes or lumps, over and above regular health practitioner consultation and/or mammography screening at the recommended interval for their risk group. Others involve attendance for consultation at a medical practice or health service and consent to examination or procedure. We focus on specific activities relating to prevention of cervical and breast cancers.

Why do some women attend to screening or engage in regular self-examination, whilst others do not? A further important question is whether there are socioeconomic covariates or identifiable predisposing or inhibiting psychological factors relating to early cancer detection behaviours. Education has been shown to be an important predictor of a number of health outcomes and interventions, and lower educational attainment may be hypothesised as a risk factor for low levels of ECDB, for example because of lack of awareness of the role of ECDB in preventing cancer.

There is a large literature reporting investigations of the role of personality type or psychological factors such as self-esteem, ‘locus of control’ (internal or external) or attributional styles and coping in a wide range of specific diseases and health-related and treatment-seeking behaviours. The Health Belief Model has been proposed since the 1950s as a predictor of preventive health behaviour15 such as participation in a screening programme.9 We therefore considered it important to investigate psychological factors as possible covariates of ECDB items.

Familial patterns have been shown to have an impact on health care use and treatment-seeking behaviour, as well as on disease susceptibility, with genetic factors contributing substantially to variability.16 We sought to identify the contributions to variation in both individual preventive behaviour and in screening compliance behaviour using a classic twin study design comparing differences of monozygotic (MZ) and dizygotic (DZ) twins reared together.17,18 We describe an exploratory study which sought to examine the genetic and environmental influences on early cancer detection behaviours for both cervical and breast cancers, and their association with potential socio-demographic and psychological covariates.

Materials and methods

Sample

Participants were members of a cohort of 1979 female twin pairs, ascertained originally in 1980–82 from the Australian National Health and Medical Research Council Twin Register,19–21 and followed up in 1988–90 when the minimum age of respondents was 25 years.22–24 The 1988 questionnaire included items on age, sex, zygosity, birth order, level of contact, education, reproductive history, psychological traits and a number of physical and psychiatric symptoms. The questionnaire replicated most of the earlier questions and added new health items for women including early cancer detection behaviours. A two-year follow-up was carried out following the 1989 survey in order to measure medium-term stability of reports over time. Identical questionnaires were mailed in 1990 to the first 500 female individual twins who responded in 1988.

Responses to the 1988 questionnaire were obtained from both members of 1504 female twin pairs (952 MZ pairs and 552 DZ pairs). Additional
twin's date of birth and date of return of the questionnaire (EPQ-R(S)). 30 The TPQ was designed to assess three higher order personality dimensions defined by Cloninger’s unified biosocial theory of personality.31 Harm Avoidance (plus four subscales of anticipatory worry and pessimism, tension about uncertainty or physical danger, shyness with strang-ers, fatiguability and asthenia), Novelty Seeking (with six dimensions of exploratory versus rigid, disorderly versus regimented, excitable/fickle versus stoic/loyal, impulsive versus reflective, dramatic/talkative versus laconic/listener, extravagant versus frugal plus three subscales of exploratory excitability versus stoic rigidity, impulsiveness versus reflection, extravagance versus reserve), and Reward Dependence (with two subscales of social sensitivity versus detachment, and persistence). The short-form EPQ-

advancing age, and because of concern that ECDBs were not sufficiently practised by women in the age groups most at risk. Age was calculated from the twin’s date of birth and date of return of the survey.

Fundamental to the twin method is the assumption that the environments of MZ co-twins are no more similar than those of DZ co-twins – or if they are, that this does not influence intrapair similarity in the variable being analysed. Environmental similarity was measured by level of contact between co-twins. Level of contact was determined from responses to the following question: ‘How often have you and your twin SEEN and CONTACTED each other during the last few years? (1) We live together, (2) Almost every day, (3) At least once a week, (4) Once or twice a month, (5) A few times a year, (6) Less often, (7) Not at all.’ Twins were asked to check one of these seven responses for seeing and contact with their co-twin for each of the previous 9 years (since the 1980–82 survey). A general contact variable for each year was created by taking the lower number (higher level of contact) of each of the SEEN and CONTACTED responses. This information was then condensed into an overall contact variable by averaging the general contact variable over the 9 years.

Highest educational level achieved was assessed on a 7 category scale: (1) Less than 7 years’ schooling, (2) 8–10 years’ schooling, (3) 11–12 years’ schooling, (4) Apprenticeship, diploma, etc. (5) Technical or Teachers’ College, (6) University first degree, (7) University post-graduate training. Categories were collapsed into four better to reflect real distinctions between levels of attainment and to increase small cell sizes at each end of the scale: (1) 1 and 2; (2) 3; (3) 4 and 5; (4) 6 and 7.

Major items included in the questionnaire were a short form 54-item version of the Tridimensional Personality Questionnaire (TPQ)29 and the short-form 48-item revised Eysenck Personality Questionnaire (EPQ-R(S)).30 The TPQ was designed to assess three higher order personality dimensions defined by Cloninger’s unified biosocial theory of personality.31 Harm Avoidance (plus four subscales of anticipatory worry and pessimism, tension about uncertainty or physical danger, shyness with strangers, fatiguability and asthenia), Novelty Seeking (with six dimensions of exploratory versus rigid, disorderly versus regimented, excitable/fickle versus stoic/loyal, impulsive versus reflective, dramatic/talkative versus laconic/listener, extravagant versus frugal plus three subscales of exploratory excitability versus stoic rigidity, impulsiveness versus reflection, extravagance versus reserve), and Reward Dependence (with two subscales of social sensitivity versus detachment, and persistence). The short-form EPQ-

Items

Early cancer detection behaviours (ECDB) Female twins were asked to answer the following questions:

1. ‘Have you ever had a (Pap) smear test for cancer of the cervix? Yes No’ (PAP)
2. ‘Do you check your own breasts for lumps or changes? No Yes, occasionally Yes, monthly’ (Breast self-examination or BSE)
3. ‘Have you ever had a breast examination by a doctor or nurse? Yes No’ (Breast examination or BE)
4. ‘Have you ever had a mammogram (breast X-ray)? Yes No’ (MAM)

When the present study was implemented mammography was used primarily for diagnostic rather than screening purposes. Variables were recoded for consistent direction.

Hypothesised covariates Variables tested for association with each ECDB item were age, level of contact between co-twins (to assess relationship with ECDB similarity, one aspect of equal environments assumption), and educational level. Age at response was included because of the increased risk of breast and cervical cancers associated with

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items concerning similarity in appearance and being mistaken by others were included to determine zygosity.19 Pairs giving inconsistent responses were recontacted for clarification. Such questionnaires have been shown to give at least 95% agreement with diagnosis based on extensive blood-typing.25–27 More recently, members of a subsample of 198 same-sex pairs from this group, who reported themselves to be MZ, were typed for 11 independent highly polymorphic markers in the course of an asthma study; no errors in our previous zygosity diagnosis were detected.28 Of 131 like-sex pairs (male and female) who reported themselves to be DZ and who had DNA available, five (38%) were concordant at the 11 loci, with a probability of monozygosity of over 0.9999. This gave a sensitivity of self-report monozygosity of 0.98 (exact 95% CI 0.94–0.99) and a specificity of 1.00 (0.97–1.00) in this sample.28

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R(S) assesses the personality dimensions of Extraversion (E); Neuroticism (N); Social Conformity or Lie (L), and Toughmindedness, or Psychoticism (P).\textsuperscript{32} Anxiety and depression were assessed by the seven sA and seven sD items from the validated Delusions–Symptoms–States–Inventory (DSSI)\textsuperscript{33,34} and 19 items from the Hopkins Symptom Check List (SCL).\textsuperscript{35,36} Ten self-esteem items\textsuperscript{37} were included. Eight items based on Pearlin and Schooler’s mastery scale were used to formulate ‘perceived control’, conceptualised as a characteristic concerning the extent to which one’s life chances were under one’s own control or that of fate.\textsuperscript{38} Ten interpersonal dependency items based on the Emotional Reliance on Another Person dimension of the Interpersonal Dependency Inventory\textsuperscript{39} were chosen to reflect need to rely on valued others.\textsuperscript{40} Thirteen ‘ways of coping’ items\textsuperscript{41,42} comprising three factors of reliance on social support, problem-focused coping and denial factors\textsuperscript{43} were also included.

We proposed that poor compliance with or lower rates of ECDBs would be associated with personality and psychological phenotypes as follows: low Neuroticism, high Psychoticism, high Lie, low Reward Dependence, low Novelty-Seeking, low Harm Avoidance, higher depression and lower anxiety scores, lower perceived control, higher denial, lower problem-focused coping and higher ‘turning to others’ coping scores.

Data summary and model-fitting

The liability to each ECDB is assumed to have an underlying normal distribution. The appropriate statistic that estimates this correlation in liability between twins is the polychoric correlation, of which the special case for a dichotomous variables is the tetrachoric correlation. To overcome scaling problems while retaining the characteristics of the original distributions, categories were created for the larger psychological scales by subtracting the number which would make ‘1’ the lowest category, then dividing the continuous score to reduce it to less than 15, the maximum number of categories allowable for calculation of polychoric correlations in PRELIS.

Matrices of correlations, and corresponding asymptotic covariance matrices, were computed separately for MZ and DZ twin pairs using PRELIS 2.12.\textsuperscript{44} Genetic models were fitted by the method of asymptotic (weighted) least squares (WLS) using LISREL 8.\textsuperscript{45} Genetic models estimating the contributions of additive genetic (a), shared or common environment (c) or nonadditive genetic influences (d) such as dominance or epistasis,\textsuperscript{46} and non-shared or specific (e) environmental effects. Genetic non-additivity and shared environment are completely confounded in data on twin pairs reared together\textsuperscript{17,47} and only one of them may be estimated.

We proceeded by systematically testing the significance of dropping parameters in turn. In addition to the likelihood ratio $\chi^2$ test (LR), the Akaike Information Criterion (AIC, measured as $\chi^2 - 2df$) was used as an additional indicator of fit. The model using the smallest number of different parameters to produce accurately the observed correlations is accepted as the best model and thus the ‘simplest’ explanation of the data (principle of parsimony). Univariate model-fitting proceeded through a systematic process of fitting of a full model containing $a$, $c$, and $e$ (ACE model), then an AE model dropping $c$, a CE model dropping $a$, and then a full ADE followed by a nested E only model with both $a$ and $d$ dropped to assess effect on fit (see Results, Table 5). For multivariate model-fitting we used a saturated Cholesky (genetic decomposition) model to explain sources of variation and covariation.\textsuperscript{48} Twin pair data analysis methods are described more fully elsewhere.\textsuperscript{48–51} Following multivariate model-fitting, parameter estimates were rotated independently using varimax rotation in SAS.\textsuperscript{52}

Results

Test–retest reliability of ECDB items

Changes in twins’ responses between 1988 and 1990, and polychoric correlations between original responses to the four ECDB questions and follow-up responses are shown in Table 1. Repeatability was highest for PAP and MAM. It is feasible that responses that were originally ‘no’ legitimately changed to ‘yes’ over the two years between the original mailing and the follow-up survey. Inconsistent answers, those that changed from ‘yes’ to ‘no’, should be equally as likely as those that are not legitimate changes from ‘no’ to ‘yes’. By setting the number of changes of ‘no’ to ‘yes’ to be equal to the number of changes of ‘yes’ to ‘no’, with the remaining number of changes of ‘no’ to ‘yes’ being deemed the legitimate changes, correlations were recomputed and coefficients increased in magnitude, particularly in the case of mammograms (see Table 1).

Covariates

Age Of the 3903 individual females who responded to the survey, 3771 (96.6%) answered all four ECDB questions. The sample was divided into age bands: 24–29, 30–39, 40–49, 50–59, 60–69 and 70+ years. Frequencies of response by age group are shown in Table 2.
Reported participation in early cancer detection behaviours differed according to age at response. Endorsement of having had a Pap smear test was lower in the youngest and oldest age groups, and was especially low in the group aged 70 and above. Rates of endorsement were very high among women in their middle years. There appear to be similar participation levels in breast self-examination across age groups although the majority of women were not performing BSE monthly and a higher number were never using it as an ECDB. BSE was much less common than BE, although for women aged 60 and over reported BE was lower than BSE. Low participation in MAM was expected as screening programmes for breast cancer were not widespread at the time of the study. Younger women had the lowest level of participation, but the relatively low rate for women aged 60 and above was noteworthy given their increased risk.

The sample was divided into two groups at the median age – the younger group aged 38 years or less and the older group aged 39 years or more – to assess differences in twin pair correlations between groups. Correlations were consistent across age groups (see Table 3). Although the pooled correlations were slightly inflated in comparison with the component correlations for each age group, we considered that the effect would not account for more than a trivial proportion of the variance, and age was not included as a covariate in model-fitting.

**Level of contact** Categories were collapsed into two categories of contact – high versus low (see Table 4). Twin pair correlations did not differ for low and high contact pairs, suggesting that although MZ co-twins are in higher overall contact, their greater similarity for each ECDC was unlikely to be caused by greater MZ co-twin contact.

### Table 1
Changes in response 1988 to 1990 and test–retest repeatability coefficients

<table>
<thead>
<tr>
<th>Response in 1990</th>
<th>PAP</th>
<th>BSE</th>
<th>BE</th>
<th>MAM</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Response in 1988</strong></td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Yes</td>
<td>402</td>
<td>5</td>
<td>367</td>
<td>31</td>
</tr>
<tr>
<td>No</td>
<td>10</td>
<td>33</td>
<td>16</td>
<td>33</td>
</tr>
<tr>
<td>Polychoric $r^a$</td>
<td>0.97</td>
<td>0.02</td>
<td>0.81</td>
<td>0.06</td>
</tr>
<tr>
<td>Polychoric $r^b$</td>
<td>0.98</td>
<td>0.02</td>
<td>0.88</td>
<td>0.04</td>
</tr>
</tbody>
</table>

$^a$PAP = ever had a Pap smear; BSE = performing breast self-examination; BE = ever had a breast examination; MAM = ever had a mammogram.

### Table 2
Frequencies of responses to the early cancer detection behaviour items in 1989 for female twins by age band

<table>
<thead>
<tr>
<th>Age band</th>
<th>n</th>
<th>%</th>
<th>%</th>
<th>%</th>
<th>%</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>24–29</td>
<td>701</td>
<td>86</td>
<td>14</td>
<td>13</td>
<td>69</td>
<td>18</td>
</tr>
<tr>
<td>30–39</td>
<td>1304</td>
<td>96</td>
<td>4</td>
<td>19</td>
<td>68</td>
<td>13</td>
</tr>
<tr>
<td>40–49</td>
<td>858</td>
<td>96</td>
<td>4</td>
<td>24</td>
<td>65</td>
<td>11</td>
</tr>
<tr>
<td>50–59</td>
<td>442</td>
<td>93</td>
<td>7</td>
<td>23</td>
<td>68</td>
<td>9</td>
</tr>
<tr>
<td>60–69</td>
<td>332</td>
<td>82</td>
<td>18</td>
<td>22</td>
<td>62</td>
<td>16</td>
</tr>
<tr>
<td>70+</td>
<td>134</td>
<td>64</td>
<td>36</td>
<td>12</td>
<td>73</td>
<td>15</td>
</tr>
<tr>
<td>Total</td>
<td>3771</td>
<td>92</td>
<td>8</td>
<td>19</td>
<td>67</td>
<td>14</td>
</tr>
</tbody>
</table>

### Table 3
Twin pair correlations (standard errors) by zygosity and age group

<table>
<thead>
<tr>
<th>Zygosity</th>
<th>Age group</th>
<th>n pairs</th>
<th>PAP</th>
<th>BSE</th>
<th>BE</th>
<th>MAM</th>
</tr>
</thead>
<tbody>
<tr>
<td>MZ</td>
<td>Younger$^a$</td>
<td>441</td>
<td>0.67 (0.09)</td>
<td>0.37 (0.07)</td>
<td>0.32 (0.08)</td>
<td>0.47 (0.10)</td>
</tr>
<tr>
<td></td>
<td>Older$^b$</td>
<td>444</td>
<td>0.68 (0.09)</td>
<td>0.37 (0.06)</td>
<td>0.45 (0.09)</td>
<td>0.46 (0.07)</td>
</tr>
<tr>
<td></td>
<td>Total</td>
<td>885</td>
<td>0.68 (0.06)</td>
<td>0.38 (0.04)</td>
<td>0.40 (0.06)</td>
<td>0.51 (0.05)</td>
</tr>
<tr>
<td>DZ</td>
<td>Younger</td>
<td>262</td>
<td>0.24 (0.19)</td>
<td>0.21 (0.09)</td>
<td>0.14 (0.11)</td>
<td>0.19 (0.17)</td>
</tr>
<tr>
<td></td>
<td>Older</td>
<td>264</td>
<td>0.14 (0.18)</td>
<td>0.14 (0.09)</td>
<td>0.13 (0.13)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td></td>
<td>Total</td>
<td>526</td>
<td>0.19 (0.13)</td>
<td>0.18 (0.06)</td>
<td>0.15 (0.06)</td>
<td>0.24 (0.09)</td>
</tr>
</tbody>
</table>

$^a$aged 38 years or less; $^b$aged 39 years or more.
Nonadditive genetic influences, although indicated in all cases resulted in a significant worsening of fit. Dropping additive genetic influences from the model (CE) in worse than the ACE model (see Table 5). Dropping shared environment offered a fit not significantly worse than the ECDB of interest were PAP ($r = 0.02$), BSE ($r = 0.002$), BE ($r = -0.02$) and MAM ($r = 0.12$) ($n = 3740$). We did not therefore include education in multivariate model-fitting.

**Psychological measures** Sample sizes without missing values for phenotypic correlations ranged from $n = 3032$ to $n = 3730$. Correlation coefficients were at best extremely modest for all scales and subscales. The strongest associations were related to PAP: the largest coefficient was with the TPQ Reward Dependence subscale of social sensitivity ($r = 0.18$), the second largest coefficient ($r = 0.16$) suggesting a small association with the TPQ Novelty Seeking dimension of ‘extravagant vs frugal’, the third largest was the inverse correlation with the shyness with strangers subscale of the TPQ Harm Avoidance dimension ($r = -0.15$). Others were negligible.

The highest correlation between any ECDB and the EPQ-R(S) Neuroticism dimension was with BSE ($r = 0.07$). Extraversion was negatively correlated at very low levels (less than 0.09) with all ECDBs. The strongest association involving self-esteem, interpersonal dependency, perceived control and ways of coping was $r = -0.10$ between BSE and problem-focused coping. We had proposed that Harm Avoidance (HA) might be related to ECDB, but found very low associations (with PAP $r = 0.07$, with BSE $r = 0.07$, with BE $r = 0.07$ and with MAM $r = 0.00$). Because the correlations were all of such low magnitude, none was included in subsequent multivariate model-fitting.

**Univariate genetic analysis**

In the case of each ECDB, the AE model without shared environment offered a fit not significantly worse than the ACE model (see Table 5). Dropping additive genetic influences from the model (CE) in all cases resulted in a significant worsening of fit. Nonadditive genetic influences, although indicated where the MZ correlation is more than twice the DZ correlation, could be dropped from the model without significantly worsening fit. The E model offers a significantly worse fit for each ECDB, indicating substantial familial aggregation for response to PAP, BSE, BE and MAM.

The heritability of liability to having had a Pap smear was the strongest at 66%, with the breast examination variables showing similar heritabilities (37% for BSE and 38% for BE). Liability to having had a mammogram had a higher genetic contribution of 51% of total variance (see Table 5).

**Multivariate genetic analysis**

The four ECDBs were variably phenotypically intercorrelated. The highest coefficients were between BE and MAM ($r = 0.57$) and BE and PAP ($r = 0.47$), where all involved a medical or nurse consultation, even though the latter two related to different cancers. It is feasible that a breast examination and mammography, although carried out by different practitioners, could relate to the same health episode. The association between BE and BSE was lower ($r = 0.26$), as was that between MAM and BSE ($r = 0.26$), although both activities related to breast cancer. Lower still was the association between PAP and BSE ($r = 0.16$), where one ECDB was consultation-related and the other a private activity, in the latter case relating to different cancers. The correlation between PAP and MAM was also modest ($r = 0.21$), although both involved consultations.

The extent of genetic and environmental covariation between the four ECDBs was assessed by fitting an atheoretical Cholesky decomposition model to polychoric correlation and asymptotic covariance matrices. As univariate analyses had suggested the AE model to be the best fitting and most parsimonious model, we fitted a double (AE) Cholesky model containing only additive genetic and specific environmental parameters. This model offered a good fit with $\chi^2_{lo} = 33.18$ ($P = 0.77$, AIC = -46.82). Estimates of contributions to variance and covariance are shown in Table 6. Parameter estimates for the four

---

### Table 4 Twin pair correlations by zygosity and level of contact of co-twins

<table>
<thead>
<tr>
<th>Zygosity</th>
<th>Level</th>
<th>n</th>
<th>PAP</th>
<th>BSE</th>
<th>BE</th>
<th>MAM</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>$r$ (SE)</td>
<td>$r$ (SE)</td>
<td>$r$ (SE)</td>
<td>$r$ (SE)</td>
</tr>
<tr>
<td>MZ</td>
<td>Love</td>
<td>256</td>
<td>0.73 (0.10)</td>
<td>0.32 (0.09)</td>
<td>0.26 (0.12)</td>
<td>0.58 (0.09)</td>
</tr>
<tr>
<td></td>
<td>High</td>
<td>627</td>
<td>0.65 (0.07)</td>
<td>0.39 (0.05)</td>
<td>0.44 (0.06)</td>
<td>0.48 (0.07)</td>
</tr>
<tr>
<td></td>
<td>Total</td>
<td>883</td>
<td>0.68 (0.06)</td>
<td>0.38 (0.04)</td>
<td>0.40 (0.06)</td>
<td>0.51 (0.05)</td>
</tr>
<tr>
<td>DZ</td>
<td>Low</td>
<td>259</td>
<td>0.22 (0.21)</td>
<td>0.22 (0.08)</td>
<td>0.15 (0.11)</td>
<td>0.23 (0.12)</td>
</tr>
<tr>
<td></td>
<td>High</td>
<td>264</td>
<td>0.16 (0.17)</td>
<td>0.13 (0.10)</td>
<td>0.16 (0.12)</td>
<td>0.26 (0.13)</td>
</tr>
<tr>
<td></td>
<td>Total</td>
<td>523</td>
<td>0.19 (0.13)</td>
<td>0.18 (0.06)</td>
<td>0.15 (0.08)</td>
<td>0.24 (0.09)</td>
</tr>
</tbody>
</table>

* The low contact group included twins who were in contact once or twice a month, a few times a year, less often or not at all; *The high contact group included twins who either lived together, contacted each other almost every day, or at least once a week.

---

*Educational level* Correlation coefficients between the four-level highest educational attainment with the ECDB of interest were PAP ($r = 0.02$), BSE ($r = 0.002$), BE ($r = -0.02$) and MAM ($r = 0.12$) ($n = 3740$). We did not therefore include education in multivariate model-fitting.

*Psychological measures* Sample sizes without missing values for phenotypic correlations ranged from $n = 3032$ to $n = 3730$. Correlation coefficients were at best extremely modest for all scales and subscales. The strongest associations were related to PAP: the largest coefficient was with the TPQ Reward Dependence subscale of social sensitivity ($r = 0.18$), the second largest coefficient ($r = 0.16$) suggesting a small association with the TPQ Novelty Seeking dimension of ‘extravagant vs frugal’, the third largest was the inverse correlation with the shyness with strangers subscale of the TPQ Harm Avoidance dimension ($r = -0.15$). Others were negligible.

The highest correlation between any ECDB and the EPQ-R(S) Neuroticism dimension was with BSE ($r = 0.07$). Extraversion was negatively correlated at very low levels (less than 0.09) with all ECDBs. The strongest association involving self-esteem, interpersonal dependency, perceived control and ways of coping was $r = -0.10$ between BSE and problem-focused coping. We had proposed that Harm Avoidance (HA) might be related to ECDB, but found very low associations (with PAP $r = 0.07$, with BSE $r = 0.07$, with BE $r = 0.07$ and with MAM $r = 0.00$). Because the correlations were all of such low magnitude, none was included in subsequent multivariate model-fitting.

*Univariate genetic analysis* In the case of each ECDB, the AE model without shared environment offered a fit not significantly worse than the ACE model (see Table 5). Dropping additive genetic influences from the model (CE) in all cases resulted in a significant worsening of fit. Nonadditive genetic influences, although indicated where the MZ correlation is more than twice the DZ correlation, could be dropped from the model without significantly worsening fit. The E model offers a significantly worse fit for each ECDB, indicating substantial familial aggregation for response to PAP, BSE, BE and MAM.

The heritability of liability to having had a Pap smear was the strongest at 66%, with the breast examination variables showing similar heritabilities (37% for BSE and 38% for BE). Liability to having had a mammogram had a higher genetic contribution of 51%of total variance (see Table 5).

*Multivariate genetic analysis* The four ECDBs were variably phenotypically intercorrelated. The highest coefficients were between BE and MAM ($r = 0.57$) and BE and PAP ($r = 0.47$), where all involved a medical or nurse consultation, even though the latter two related to different cancers. It is feasible that a breast examination and mammography, although carried out by different practitioners, could relate to the same health episode. The association between BE and BSE was lower ($r = 0.26$), as was that between MAM and BSE ($r = 0.26$), although both activities related to breast cancer. Lower still was the association between PAP and BSE ($r = 0.16$), where one ECDB was consultation-related and the other a private activity, in the latter case relating to different cancers. The correlation between PAP and MAM was also modest ($r = 0.21$), although both involved consultations.

The extent of genetic and environmental covariation between the four ECDBs was assessed by fitting an atheoretical Cholesky decomposition model to polychoric correlation and asymptotic covariance matrices. As univariate analyses had suggested the AE model to be the best fitting and most parsimonious model, we fitted a double (AE) Cholesky model containing only additive genetic and specific environmental parameters. This model offered a good fit with $\chi^2_{lo} = 33.18$ ($P = 0.77$, AIC = -46.82). Estimates of contributions to variance and covariance are shown in Table 6. Parameter estimates for the four
additive genetic and the four individual environmental factors were rotated independently using varimax rotation in SAS (see Table 7).

Genetic covariation between the ECDBs was minimal, with the exception of a loading from BE on the first genetic factor (a PAP factor), suggesting some common genetic influence. Environmental covariation was low in all cases. Results suggested that no common factor, either environmental or genetic, underlies the four behaviours and that fitting a common pathway model to the data was not warranted.

Table 5  Twin pair polychoric correlations and proportions of variance from univariate genetic model-fitting for early cancer detection behaviours

<table>
<thead>
<tr>
<th>Proportions of variance</th>
<th>ECDB</th>
<th>Model</th>
<th>$a^2$</th>
<th>$c^2$</th>
<th>$e^2$</th>
<th>$d^2$</th>
<th>$\chi^2$</th>
<th>df</th>
<th>P</th>
<th>AIC</th>
</tr>
</thead>
<tbody>
<tr>
<td>PAP</td>
<td></td>
<td>ACE</td>
<td>0.66</td>
<td>0.00</td>
<td>0.34</td>
<td>-</td>
<td>1.29</td>
<td>0</td>
<td>1.00</td>
<td>1.29</td>
</tr>
<tr>
<td>$r_{MZ} = 0.68^a$</td>
<td></td>
<td>AE</td>
<td>0.66</td>
<td>-</td>
<td>0.34</td>
<td>-</td>
<td>1.29</td>
<td>1</td>
<td>0.26</td>
<td>-0.71</td>
</tr>
<tr>
<td>$r_{DZ} = 0.19^a$</td>
<td></td>
<td>CE</td>
<td>-</td>
<td>0.59</td>
<td>0.41</td>
<td>-</td>
<td>11.70</td>
<td>1</td>
<td>0.00</td>
<td>9.70</td>
</tr>
<tr>
<td>ADE</td>
<td>0.07</td>
<td>0.32</td>
<td>0.61</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.00</td>
<td>0</td>
<td>1.00</td>
<td>0.00</td>
</tr>
<tr>
<td>E</td>
<td>-</td>
<td>1.00</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>135.79</td>
<td>2</td>
<td>0.00</td>
<td>131.79</td>
<td></td>
</tr>
<tr>
<td>BSE</td>
<td></td>
<td>ACE</td>
<td>0.37</td>
<td>0.00</td>
<td>0.63</td>
<td>-</td>
<td>0.01</td>
<td>0</td>
<td>1.00</td>
<td>0.01</td>
</tr>
<tr>
<td>$r_{MZ} = 0.37$</td>
<td></td>
<td>CE</td>
<td>-</td>
<td>0.30</td>
<td>0.70</td>
<td>-</td>
<td>6.49</td>
<td>1</td>
<td>0.01</td>
<td>4.49</td>
</tr>
<tr>
<td>$r_{DZ} = 0.18$</td>
<td></td>
<td>E</td>
<td>-</td>
<td>-</td>
<td>0.63</td>
<td>0.02</td>
<td>-</td>
<td>0</td>
<td>1.00</td>
<td>0.00</td>
</tr>
<tr>
<td>BE</td>
<td>0.39</td>
<td>0.61</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>76.69</td>
<td>2</td>
<td>0.00</td>
<td>72.69</td>
</tr>
<tr>
<td>$r_{MZ} = 0.40$</td>
<td></td>
<td>AE</td>
<td>0.39</td>
<td>0.00</td>
<td>0.63</td>
<td>-</td>
<td>0.31</td>
<td>0</td>
<td>1.00</td>
<td>0.31</td>
</tr>
<tr>
<td>$r_{DZ} = 0.15$</td>
<td></td>
<td>CE</td>
<td>-</td>
<td>0.31</td>
<td>0.69</td>
<td>-</td>
<td>6.15</td>
<td>1</td>
<td>0.01</td>
<td>4.15</td>
</tr>
<tr>
<td>$r_{DZ} = 0.24$</td>
<td></td>
<td>E</td>
<td>-</td>
<td>-</td>
<td>0.61</td>
<td>0.19</td>
<td>-</td>
<td>0</td>
<td>1.00</td>
<td>0.00</td>
</tr>
<tr>
<td>MAM</td>
<td></td>
<td>ACE</td>
<td>0.51</td>
<td>0.00</td>
<td>0.49</td>
<td>-</td>
<td>0.02</td>
<td>0</td>
<td>1.00</td>
<td>0.02</td>
</tr>
<tr>
<td>$r_{MZ} = 0.51$</td>
<td></td>
<td>CE</td>
<td>-</td>
<td>0.44</td>
<td>0.56</td>
<td>-</td>
<td>6.74</td>
<td>1</td>
<td>0.01</td>
<td>4.74</td>
</tr>
<tr>
<td>$r_{DZ} = 0.24$</td>
<td></td>
<td>E</td>
<td>-</td>
<td>-</td>
<td>0.49</td>
<td>0.05</td>
<td>0.00</td>
<td>0</td>
<td>1.00</td>
<td>0.00</td>
</tr>
<tr>
<td>E</td>
<td>0.01</td>
<td>0.07</td>
<td>0.19</td>
<td>0.24</td>
<td>0.51</td>
<td>0.03</td>
<td>0.01</td>
<td>0</td>
<td>0.35</td>
<td>0.49</td>
</tr>
</tbody>
</table>

Best-fitting, most parsimonious models are in bold; $a^2$ = additive genetic influences; $c^2$ = common environment; $e^2$ = unique environmental influences; $d^2$ = non-additive genetic influences; $^aMZ$ n = 885 pairs; $^bDZ$ n = 523 pairs.

Table 6  Multivariate Cholesky decomposition of variance and covariance for four ECDB items (AE model)

<table>
<thead>
<tr>
<th>Proportions of variance</th>
<th>ECDB</th>
<th>Genetic factors</th>
<th>A1</th>
<th>A2</th>
<th>A3</th>
<th>A4</th>
<th>Total</th>
<th>Unique environmental factors</th>
<th>E1</th>
<th>E2</th>
<th>E3</th>
<th>E4</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>PAP</td>
<td></td>
<td>A1</td>
<td>0.66</td>
<td></td>
<td></td>
<td></td>
<td>0.34</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.34</td>
</tr>
<tr>
<td>BSE</td>
<td>0.02</td>
<td>0.36</td>
<td></td>
<td>0.38</td>
<td></td>
<td></td>
<td>0.62</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.62</td>
</tr>
<tr>
<td>BE</td>
<td>0.18</td>
<td>0.03</td>
<td>0.18</td>
<td></td>
<td></td>
<td></td>
<td>0.08</td>
<td>0.01</td>
<td>0.52</td>
<td></td>
<td></td>
<td></td>
<td>0.61</td>
</tr>
<tr>
<td>MAM</td>
<td>0.01</td>
<td>0.07</td>
<td>0.19</td>
<td>0.24</td>
<td></td>
<td></td>
<td>0.51</td>
<td>0.03</td>
<td>0.01</td>
<td>0.10</td>
<td></td>
<td>0.35</td>
<td>0.49</td>
</tr>
</tbody>
</table>

Table 7  Genetic and environmental parameter estimates following independent varimax rotation of factor estimates, with factors reordered

<table>
<thead>
<tr>
<th>Parameter estimates</th>
<th>ECDB</th>
<th>Genetic factors</th>
<th>A1</th>
<th>A2</th>
<th>A3</th>
<th>A4</th>
<th>E1</th>
<th>E2</th>
<th>E3</th>
<th>E4</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>PAP</td>
<td>0.79</td>
<td>0.06</td>
<td>0.17</td>
<td></td>
<td></td>
<td>0.04</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.38</td>
</tr>
<tr>
<td>BSE</td>
<td>0.05</td>
<td>0.59</td>
<td>0.07</td>
<td></td>
<td></td>
<td>0.10</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.03</td>
</tr>
<tr>
<td>BE</td>
<td>0.31</td>
<td>0.12</td>
<td>0.45</td>
<td></td>
<td></td>
<td>0.26</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.14</td>
</tr>
<tr>
<td>MAM</td>
<td>0.05</td>
<td>0.14</td>
<td>0.15</td>
<td></td>
<td></td>
<td>0.69</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.07</td>
</tr>
</tbody>
</table>
Discussion

The analysis of causes of individual differences in early cancer detection behaviours has produced evidence of specific genetic influences on the liability to each of the four behaviours: Pap smear, breast self-examination, breast examination and mammogram. Despite modest to strong phenotypic intercorrelations between the variables, the only genetic covariation identified was between PAP and BE. It is plausible, given the finding that genetic influences may influence health treatment-seeking behaviour, that there is a predisposition towards either initiating or complying with available health care directed at prevention by early diagnosis. We had no other measures of use of primary health care services with which to compare these findings. These behaviours appear to be distinct in aetiology from the private behaviour of breast self-examination.

The strongest genetic influences were identified for liability to ever having had a Pap smear. A plausible explanation of this finding is difficult, and was not suggested by any of the covariates tested. If it were the age of first Pap smear, possibly age of onset of sexual activity might be suggested. It is possible that a sexual activity factor might be involved, such as number of partners influencing perceived risk. A Pap smear involves willingness to undergo a vaginal examination by a doctor (or less frequently a nurse) and there can be some discomfort or, occasionally, pain. Factors may include fear of the process, being too busy caring for family members to think about their own health, or being embarrassed, especially if the doctor is male. These factors may not be assessed by the psychological measures included in the questionnaire. It is interesting that in one Australian study almost a quarter of the women screened had their Pap smear taken in the context of their use of either contraception or hormones, or in connection with either pregnancy or a postpartum visit. Oral contraceptive use has been associated with the decision to have a cervical smear test in other studies. Genetic influences are plausible on these factors as possible covariates, and may justify further investigation. Mammography in our study would have been primarily used in response to signs or symptoms, so its meaning cannot be readily extended to asymptomatic women’s predisposition to engage in mammographic screening. The genetic influences identified may relate in part at least to breast symptoms. Any effect of (knowledge of) family history would be identified as a shared environmental influence.

Despite a noticeably higher frequency of negative response by women in the older age group to the questions on Pap smear and breast examination, age was not relevant as a covariate for genetic analyses. None of the other hypothesized covariates – level of contact, educational level and personality scores for harm avoidance, novelty seeking, reward dependence, neuroticism, anxiety, depression, self esteem, perceived control interpersonal dependency and ways of coping – was found to be significant. Clearly there is a need for further investigation to find measures that do have explanatory relevance.

One might have expected genetic influences to be stronger on a private behaviour like BSE, but the proper technique does need to be taught to women by a health practitioner in the first instance and practitioner reinforcement of the practice may well be important for its continuation. In addition to women’s own knowledge and motivation, health service and practitioner factors may constitute important environmental influences affecting each ECDB. Continuity of care from a general practitioner and increased length of attendance at a practice has been linked to better practice of preventive health behaviours such as breast self-examination and having regular Pap smears. There was no evidence from our data that environmental factors were the same for any two behaviours, however. Environmental influences were less important for having had a Pap smear than the other behaviours, however. Environmental influences were less important for having had a Pap smear than the other behaviours, however. Environmental influences were less important for having had a Pap smear than the other behaviours, however. Environmental influences were less important for having had a Pap smear than the other behaviours, however. Environmental influences were less important for having had a Pap smear than the other behaviours, however. Environmental influences were less important for having had a Pap smear than the other behaviours, however. Environmental influences were less important for having had a Pap smear than the other behaviours, however. Environmental influences were less important for having had a Pap smear than the other behaviours, however. Environmental influences were less important for having had a Pap smear than the other behaviours, however. Environmental influences were less important for having had a Pap smear than the other behaviours, however. Environmental influences were less important for having had a Pap smear than the other behaviours, however. Environmental influences were less important for having had a Pap smear than the other behaviours, however.
Our study was conducted before the introduction of national policies and organised programmes with clear and consistent guidelines for screening onset and rescreening intervals. This was an advantage in that twins’ behaviours were less likely to have been influenced by the same media messages. Nevertheless, the messages are by no means yet universally recognised. Following the introduction of the national breast screening programme, some women were still uncertain about why screening is important and about recommended age for commencement of mammography and intervals for rescreening. On the other hand, if there is indeed a more uniform information environment and consistent practitioner guidelines, further study of these female twin pairs may be even more informative in identifying individual factors predisposing women to preventive health behaviours now that Australia is some years down the track following the introduction of national screening programmes.

Acknowledgements

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