Sir,

With much interest we read the paper by Antoniou and colleagues (British Journal of Cancer, October 18, p 1580). These researchers described the development of a genetic model for familial breast cancer, named BOADICEA, which takes into account the simultaneous effects of BRCA1, BRCA2 and other genes. There is a considerable need for such predictions for purposes of management of women referred to genetic clinics and for the considerations of eligibility of any prophylactic interventions in both the clinical and research settings.

The BOADICEA model requires a computer program for risk estimation, which is not easy to use in clinical practice. Moreover, such a model is mostly hard to understand for clinicians. Understandable risk estimation should, however, be available to clinicians, so that they know what they are actually doing while estimating disease risks. Our concern about this aspect of the BOADICEA model was illustrated in Figure 3. The authors presented a woman aged 40 years and unaffected with cancer. Her mother and maternal aunt developed ovarian cancer and two other maternal aunts developed ovarian cancer before the age of 50 years. This family is a good example of a typical hereditary breast and ovarian cancer (HBOC) family. The risk for breast cancer in these families is increased because of the combination of both breast and ovarian cancers in the family, which highly points to BRCA1 mutations. The BRCA1 mutation probability as predicted by the BOADICEA model is high, 40.9%. The model then predicted her risk of breast cancer at only 13%. For a clinician, this risk estimation is hard to understand and feels illogical. It seems that in the risk prediction by Antoniou et al (2004) the presence of ovarian cancer is not important for the estimation of the breast cancer risk for this woman. The authors did not discuss this point.

Recently, Jonker et al (2003) have published a genetic model to predict someone’s breast cancer risk based on the family history of breast and ovarian cancer. This model can be considered as an extension of the Claus model combined with the BRCAPRO model (Claus et al, 1991; Parmigiani et al, 1998). In the Jonker et al model, the familial clustering of breast and ovarian cancer is explained by three genes, BRCA1, BRCA2 and a hypothetical third gene BRCAu. This third gene was modelled to explain all familial clustering of breast cancer unaccounted for by the BRCA1 and BRCA2 genes. The model parameters were estimated using published estimates of population incidence and relative risks.

As well as the BOADICEA model, the Jonker et al model is not easy to use in clinical practice. For this reason, we extended the easy-to-use Claus tables into the ‘Claus plus method’ based on the Jonker model (Claus et al, 1994; Van Asperen et al, 2004). Our method uses the Claus tables, but also incorporates information on the presence of ovarian cancer, bilateral breast cancer and whether there are more than two affected relatives. The formula we obtained simplifies risk estimation in familial breast cancer:

\[
0.08 + 0.40 \times \text{Claus Table} + 0.07 \times \text{ovarian cancer} + 0.08 \times \text{bilateral breast cancer} + 0.07 \times \text{multiple cases}.
\]

The formula starts with an intercept of 0.08. This is the population risk for breast cancer and the basis for further risk estimation. The value of the Claus table should be multiplied by 0.4. The formula subsequently includes the information on ovarian cancer, bilateral breast cancer and more than two affected relatives. These characteristics in the formula are one or zero. This new method might offer a good alternative for breast cancer risk estimation in clinical practice. The ‘Claus plus method’ is an easy applicable method for hand-written pedigrees and at the moment it is widely used in the Dutch cancer clinics. Based on the ‘Claus plus method’, the predicted risk of breast cancer of the 40-year-old woman in Figure 3 in the Antoniou paper was 31%. Although we are still working on the validation of our method, this risk figure is more in agreement with the risks as observed in typical HBOC families.

The authors have mentioned several improvements to be made for their multi-purpose BOADICEA model like genotype-specific incidence rates, risk for other cancers and allele frequencies. Besides this, we would like to recommend paying attention to the applicability for clinical practice. To attain full development for the BOADICEA model, the authors should be in close contact with the ultimate users in clinical practice.
_reply:

Sir,

The BOADICEA model of genetic susceptibility to breast and ovarian cancer was developed using complex segregation analysis of breast and ovarian cancer (Antoniou et al, 2002, 2004). We agree with van Asperen et al that as it stands, the model is not easy to use in clinical practice. However, the model is currently being implemented in web-based software that will provide a user-friendly tool for clinical geneticists and oncologists. We disagree however with the premise that the model is particularly hard to understand. The BOADICEA model incorporates the effects of BRCA1, BRCA2 and other genes. Although it has more risk parameters than some other models, it is conceptually quite similar to the Claus et al (1991) or BRCAPRO models (Parmigiani et al, 1998), or the model proposed by Jonker et al (2003). The major difference is the incorporation of a polygenic component to explain familial aggregation of breast cancer not attributable to BRCA1 and BRCA2.

van Asperen et al question the low breast cancer risk (13% by age 70) estimated for the index woman in Figure 3 of our paper. We agree that this estimate is probably anomalously low, due to imprecision in the BRCA1 and BRCA2 penetrance estimates we used. The average risk of breast cancer in BRCA1 mutation carriers in the first version of the BOADICEA model was estimated to be 35% by age 70 (Antoniou et al, 2002), which is much lower than the estimates used in the Jonker et al (2003). However, the BRCA1 and BRCA2 incidence rates used in the first version of BOADICEA were based on relatively small numbers of BRCA1 and BRCA2 mutation positive families (62 in total) and may therefore be imprecise.

To improve the risk prediction, we have recently refitted the BOADICEA model using additional data from two UK population-based studies of breast and ovarian cancer (Antoniou et al, 2003). The updated data set includes more than 500 BRCA1 and BRCA2 mutation positive families, and therefore the incidence rates are estimated more reliably (manuscript in preparation). In the updated version, the average risk of breast cancer in BRCA1 mutation carriers by age 70 varies between 50 and 59% depending on the year of birth. Applying the latest version of the model to the family in Figure 2 of Antoniou et al (2004), the 40-year-old woman is predicted to carry a BRCA1 mutation with probability 41% and a BRCA2 mutation with a probability 1% (very similar to the previous estimates). However, her predicted risk of developing breast cancer by age 70 is now higher, 28%, perhaps closer to the expectations of van Asperen and co-workers.

van Asperen et al question the fact that the presence of ovarian cancer in the family does not affect the breast risk. It is a feature common to BOADICEA and all the other risk prediction models, however, that the risks of breast and ovarian cancer in a family are assumed to be independent given the BRCA1 and BRCA2 genotypes. Thus, the presence of ovarian cancer in the family only affects the breast cancer risk in so far as it affects the BRCA1 and BRCA2 carrier probabilities.

We agree that the regression model of van Asperen et al (2004) can be more easily used in clinical practice. However, this model cannot deal with the complex family histories seen in genetic
clinics. Moreover, the model was derived by linear regression of the independent variables on the predictions given the Jonker et al (2003) model. Therefore, the validity of the regression formula critically depends on an as yet (to our knowledge) unvalidated model. In any event, evaluation of the accuracy of any model in predicting the correct carrier and cancer risks should be based on validation studies in independent series and not on the basis of individual families.

REFERENCES


