Familial aggregation of breast/ovarian cancer: age of onset along subsequent generations in Brazil

Agregação familiar de câncer de mama e ovário: idade de manifestação em gerações subseqüentes no Brasil

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Abstract Antecedents of familial aggregation of breast and ovarian cancer are observed in only 5-8% of all breast cancer cases. Nevertheless, this variable displays one of the highest risk ratios associated to breast cancer outcome. Despite recent identification of genetic mutations associated with familial aggregation of these tumors, mainly at BRCA1 and BRCA2 genes, knowledge on the interaction between environmental agents in these families remains quite unclear. In this paper we ascertained the correlation among ages of the onset of breast/ovarian cancer in 260 Brazilian families with those cancer aggregation. Further we estimated the median age of the onset of breast cancer among four generations. We observed that the higher the number of family cancer cases, the highest is the correlation of ages for the onset of breast cancer. We also observed an 8-10 year decline in the mean age-of-onset of breast/ovarian cancer from one generation to another in the studied families. If these results could be confirmed elsewhere, we believe that the hypothesis of interaction between environmental risks factors in families indeed showing breast/ovarian cancer aggregation is reinforced.

Key words Neoplasm; Breast; Ovary, Ovarian; Genetics

Resumo A presença de antecedentes de agregação familiar de câncer de mama e ovário é observada em apenas 5% a 8% de todos os casos de câncer de mama. Esta variável, entretanto, é uma das que apresenta maior razão de riscos para o desfecho câncer de mama. Apesar da identificação recente de mutações genéticas associadas com a agregação familiar destes tumores, sobre tudo nos genes BRCA1 e BRCA2, o conhecimento sobre as interações entre fatores ambientais e genéticos nessas famílias permanece relativamente obscuro. Neste trabalho, determinamos a correlação das idades de início de câncer de mama e ovário em 260 famílias com agrupamento de tais tumores. Posteriormente, foi estimada a idade média de início destes tumores ao longo de quatro gerações. Observou-se que quanto maior o número de casos de câncer na família, maior o coeficiente de correlação entre as idades de início destes cânceres. Foi também constatado um declínio de oito a dez anos na idade média de início do câncer nessas famílias ao longo de três gerações. Os autores acreditam que, se esses resultados forem confirmados por outras investigações, reforça-se a hipótese de uma interação entre fatores ambientais e genéticos no desenvolvimento dos tumores de mama e ovário em famílias que já apresentam uma forte agregação desses tumores.

Palavras-chave Neoplasia; Mama; Ovário; Genética
Introduction

Breast cancer has been a leading cause of cancer among women in developed countries, showing an increasing incidence and stable mortality during the last decades (Garfinkel et al., 1994). Among the known risk factors associated to that disease, familial aggregation of female breast cancer has been identified as an important condition yielding to increased risk of developing that tumor. Antecedents of first degree relatives affected with breast cancer ranges from 8% to 30% among cases with such history (Anderson & Badzioch, 1985; Dawson & Thompson, 1989; Marcus et al., 1996). Indeed, risk ratios as high as 5 has been mentioned among women with previous cases of breast cancer among close relatives, such as a mother or a sister, as compared with women without family antecedents of that neoplasm (Kelsey, 1993). For instance, if just a woman’s mother has had breast cancer, her relative risk to develop the same cancer is 1.8; if one sister has had the disease, the risk is quite higher, 2.5; moreover, if both mother and sister had developed breast cancer, the risk jumps to 5.6 (Bain et al., 1980). The reasons involved in this process remained quite unclear until the last few years when knowledge about the genetics of breast cancer experienced a remarkable improvement.

Familial breast cancer is now known to develop in multiple family relatives among subsequent generations, very often bilateral, and at relatively young ages. Up to now, some genes were identified as having an important role in familial aggregation of breast cancer. At first, the gene named BRCA1 (for breast cancer) located in chromosome 17 had been implicated in cancer expression of families displaying female breast and ovarian cancer (Hall et al., 1990). More than eighty mutations in this gene were described, despite the fact that the importance of most of them remains unknown. Another gene involved in familial aggregation of breast cancer is BRCA2 located at chromosome 13, and mainly associated with cases of male breast cancer and ovarian cancer in the same families. The existence of a third gene, still unidentified, yet named BRCA3, was discussed taking into account observation of family cases of breast and ovarian cancer without showing mutations either on BRCA1 or BRCA2. As a whole, BRCA1 and BRCA2 (or even BRCA3 if further identified) seems to play the most important role in familial aggregation of breast and ovarian cancer cases, while p53 gene seems to be more involved in the expression of sporadic breast cancer cases (Elledge & MacGuire, 1994; Ozbun & Butel, 1995). Nevertheless, p53 gene was also described in Li-Fraumeni families (Malkin et al., 1990).

The manner in which genetic and environmental risk factors act during carcinogenesis either on familial aggregation and sporadic breast cancer remains unclear. Therefore, efforts have been made trying to improve understanding of their possible interaction by describing patterns of disease spreading among populations. Early menarche, late menopause, declining parity and consumption of oral contraceptives during adolescence before a full-term pregnancy, were also pointed out as variables possibly associated with breast cancer related to a pattern of high exposure to estrogens during the reproductive years (Kelsey, 1993; Newcomb et al., 1996). Nevertheless, these variables seems to act differently if cases are sporadic or family aggregated (Colditz et al., 1996).

Among variables associated to breast cancer, age is perhaps one of the best documented in literature. Breast cancer in different populations show quite similar distributions by age, with sharp increases on incidence rates between 25 and 45 yr. old, followed by a slower increasing speed on incidence after that age. Indeed, this observation has been pointed out as a strong indicator of hormonal involvement in breast cancer development. Therefore, breast cancer is mainly a post-menopausal disease, affecting around 4% of women below 35 yrs. old (Chung et al., 1996) and 7% below 40 yrs. old (Winchester, 1996). The majority of cases are diagnosed between 45 and 55 yrs. old in the USA, and only 20% among women 75 yrs. or older (Busch et al., 1996). One of the most interesting findings recently mentioned in the literature about age distribution of breast cancer is the decline in the age of onset among breast cancer cases observed during the last decades.

Different studies have shown that age of onset among families with several cases of breast cancer usually start at younger ages than observed among sporadic cases in the general population (Dotto, 1985). Nevertheless, the manner in which cases from families with antecedents of breast and ovarian cancer behavior has been poorly documented in the literature with regard to age distribution among successive generations.

In this paper we will present data on the age of onset of breast/ovarian cancer among Brazilian families showing several cases of both tumors identified among different generations.
Implications of the observed trends will be further commented on with consideration to available knowledge in the literature.

Methods

Since 1994, the Brazilian National DNA Bank Program (BNDBP) has been gathering blood samples and epidemiological information from families showing aggregation of breast and ovarian cancer in different cities. At first Rio de Janeiro, São Paulo and Porto Alegre were included, with this list increasing with time. Blood samples have been used to identify genetic mutations, which will be further linked to epidemiological data aiming to analyze possible patterns of their distribution among these families. In this paper, we will present epidemiological information on the distribution of age for the onset among cases of breast/ovarian cancer identified among families with 3 or more breast/ovarian cancer cases in the three aforementioned cities.

At first, Pearson’s correlation coefficients between ages of onset of breast/ovarian cancer among enrolled women in BNDBP was carried out. Furthermore, distribution on the ages of the onset of breast/ovarian cancer was ascertained among subsequent generations in the same families. These ages were determined towards the median age of onset in each generation for each family, as a measure to guarantee the same probability for all families to be represented, regardless of the number of affected women in each family.

Results and discussion

Among enrolled families in BNDBP, 260 have two or more cases of breast/ovarian cancer in Rio de Janeiro and São Paulo, 64 of them with 4 or more cases. Median age at diagnosis among enrolled cases with 4 or more family cases was 52 yr. and 48 yr. among their close relatives.

As seen in Table 1, a high correlation of age at the onset between close relatives of cancerous enrolled women was observed: r = 0.95 (0.81-0.99), F = 81.2, p < 0.005. Correlation of age at the onset among close relatives of non-cancerous enrolled women with multiple family cases (4 or more) was also high, r = 0.79 (0.59-0.80), F = 84.0, p < 0.005. Among non-cancerous enrolled women with 3 or less family cases, the correlation coefficient was moderate, r = 0.43 (0.33-0.53), F = 59.3, p < 0.005.

These results suggest a strong correlation between ages at onset of breast and ovarian cancer among relatives from families with past family antecedents of these tumor sites. Moreover, the higher the number of family cases, the higher correlation was observed between ages of breast and ovarian cancer onset. Nevertheless, these results are not informative on directionality or timing about ages of onset, and they just indicate similarity between these ages among all studied families. In other words, they just reveal an important closeness or similarity between age groups in which cases were diagnosed among families with past antecedents of breast/ovarian cancer.

To analyze time trends on the age of onset of breast/ovarian cancer, Table 2 is more useful: when distribution of ages of the onset of breast and ovarian cancer was carried out along subsequent generations, a decline on the mean age of onset was observed: 53.4 yr. in the first generation of affected women, 43.3 yr. in the second, 34.6 yr. in the third and 30.5 yr. in the fourth (F = 30.85; p < 0.0001). These results – obtained towards several intra-family comparisons along subsequent generations of close relatives –, suggest a marked decline among ages in which these tumors were diagnosed, i.e., a 8-10 years decline between two subsequent generations considering the first three ones.

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<td>Distribution of cases of breast/ovarian cancer among enrolled families, Brazilian National DNA Bank Program.</td>
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These results should be interpreted cautiously, because they may be biased in the sense that women in the first generation had experienced a higher life span than those in the third or the fourth generation, who still are in their 30's or 40's. Therefore, as their follow up goes on, cases at older ages will certainly be diagnosed, increasing the mean age of onset in the respective third and fourth family generation. Moreover, mammography access may also allow breast cancer detection at earlier ages, changing the mean age-of-onset of breast cancer among different generations. Henceforth, we believe that a continuous follow up of these women should yield to closer mean ages at the onset of breast/ovarian cancer than the time interval now observed (8-10 years between subsequent generations). Nevertheless, we also consider these intervals so large, that it seems quite improbable that lifelong correction of ages at the onset, brought by a continuous follow up, could eliminate the current differences observed among women in subsequent generations.

Consequently, we believe that present results may indeed point out a true trend towards a decline in the age of the onset of breast/ovarian cancer diagnosed among women from families with a strong pattern of these tumors aggregation. Therefore, a comprehensive analyses taking into account the role of environmental risk factors to breast and ovarian cancer among families showing this tumor aggregation pattern in contemporary societies should be explored. Unfortunately, the majority of epidemiological studies analyzing family aggregation of breast cancer have still been just oriented towards the important genetic aspects involved in this process, and not considering the probable role of environmental risk factors in their expression.

Conclusions

A sample of Brazilian families showing a pattern of breast and ovarian cancer aggregation was studied. A statistically significant 8 to 10 yr. decline on the mean age of the onset of breast/ovarian cancer among their close relatives was observed along three subsequent generations.

References


