ABSTRACT

This literature review aims to inform and assist physicians and other health professionals in managing all information related to hereditary breast cancer, which is in constant and rapid growth, allowing for improvement in patient care and assistance. In addition, we seek to better identify which patients are eligible for the clinical criteria of association with risk of hereditary breast cancer, based on international recommendations and highlighting the main high and moderate penetrance genes that make up the multigenic panels for germline investigation in breast cancer, as well as the possibilities of clinical management that must be considered when complex decisions are required in clinical practice. Nowadays, there is more interest in population screening, in a greater supply of genetic tests, more genes included in multigene panels — allowing the search for genetic counseling —, apart from the need for clinical-decision support.

KEYWORDS: hereditary breast and ovarian cancer syndrome; screening; genetic counseling; genetic testing; treatments; risk reduction.

LITERATURE REVIEW

Breast cancer and hereditary predisposition

Breast cancer is the most common malignant neoplasm whose mortality rates are the highest among the females worldwide. In Brazil, 66,280 new cases were estimated per year for the triennium 2020–2022 (43.74 cases per 100,000 women)1,2. Although there are several risk factors, breast cancer is associated with environmental, reproductive, genetic and lifestyle factors; family history is considered an important etiological factor3.

The hereditary factor is a cause identified in 10–15% of breast cancer cases, and is associated with the Hereditary Breast and Ovarian Cancer Syndrome (HBOC), Li-Fraumeni Syndrome, Cowden syndrome, Peutz-Jeghers syndrome and hereditary diffuse gastric cancer in which lobular breast carcinoma may manifest in women4–6.

The hereditary predisposition is most commonly seen in individuals and families with some clinical features such as: diagnosis of breast cancer in patients ≤45 years of age; breast cancer in men; personal and family history of ovarian, pancreas, bowel, endometrial, and prostate cancers at a younger age; and Ashkenazi Jewish origin7.

Pathogenic variants

HBOC is mainly related to pathogenic variants in the BRCA1 and BRCA2 genes, which consist of germline mutations and account for almost 30% of all cases of hereditary breast cancer8,9. Other genes also have pathogenic variants associated with increased risk for hereditary breast cancer, such as TP53, CHEK2, ATM, STK11, PALB2, PTEN, among others, which demonstrates the complex genetic involvement when it comes to predisposition to this disease10.

Over a lifetime, the presence of a pathogenic variant in BRCA1 or BRCA2 can increase the risk of breast cancer by up to 85%. For ovarian cancer, estimates reach 46% when the BRCA1 gene is involved and 20% when the BRCA2 gene is involved4,11,12. The BRCA1 and BRCA2 genes, identified in the 1990s, are involved in the activation of DNA repair in response to cellular stress, playing crucial roles in chromatin remodeling, transcriptional control and cell cycle regulation, with tumor suppressor effects primarily attributed to cell cycle checkpoints and DNA repair12–15. Some mutations are more common in individuals from specific ethnic or geographic groups. This is due to the presence of initiating mutations which probably arose several generations ago in
this population. In Ashkenazi Jews (descendants of Central and Eastern Europe), three specific initiating mutations were identified: 185delAG and 5382insC in the *BRCA1* gene, and 6174delT in the *BRCA2* gene, and the same family group may have all three mutations. These variants are present in 2% of individuals in this group of women and are responsible for approximately 50% of early-onset breast cancer cases.

Another founder mutation identified in Portuguese Caucasian families with cases of breast cancer is the insertion Aluc.156_157insALU in exon 3 of the *BRCA2* gene, which promotes DNA rearrangements, altering the nucleotide sequence. The Brazilian ethnic composition also makes room for founder mutations in our population.

Another tumor suppressor gene called *TP53*, associated with the Li-Fraumeni syndrome, leads to increased risk for multiple tumors, including osteosarcoma, bowel cancer, adrenocortical carcinoma, leukemia, lymphoma, and brain cancer in addition to breast cancer. This syndrome has an interesting peculiarity in patients diagnosed in the South and Southeast regions of Brazil: the founder mutation p.R337H has a prevalence of 0.3% due to the founder effect related to the movement of drovers in Brazilian territory.

The *PTEN* gene, responsible for cell cycle control, is associated with the Cowden syndrome and usually causes malignant tumors in the thyroid, breast and endometrium. Women with this syndrome have 25% to 50% risk of developing breast cancer, while the risk of endometrial carcinoma can reach 10%.

Pathogenic variants involving the *STK11* gene are associated with the Peutz-Jeghers syndrome, which increases the lifetime risk of breast cancer in women by up to 50%. Furthermore, genes involved in the pathways of DNA double-strand break (DSB) such as *CHEK2, RAD51, BRIP1* and *PALB2*, may also be associated with hereditary breast cancer predisposition.

Most cases of breast cancers are invasive and the prognosis depends on the stage of the disease at the time of diagnosis. In general, in developing countries, diagnoses occur at advanced stages, which is mainly due to the deficiency in promoting early detection. In non-menopausal women, breast cancer represents a biologically more aggressive disease, with frequent adverse histopathological features and worse prognosis when compared to women over 50 years of age.

Hereditary breast cancer with *BRCA1* mutation often results in triple-negative breast cancers — approximately 80% of *BRCA1* mutation cases. Histological characterization of tumors with *BRCA1* germline mutations suggests high histological grade, atypical medullary features, high proliferation rates, inflammatory infiltrates, and invasive borders. On the other hand, *BRCA2* mutation are related to tumors with a higher risk of contralateral breast cancer and estrogen receptor positivity in most cases.

### Screening for hereditary breast cancer

Although physical examination is important to establish doctor-patient relationships and to evaluate symptomatic patients, it plays a less important role in breast cancer screening when compared to imaging methods such as mammography, magnetic resonance imaging (MRI) and ultrasonography, since it has low sensitivity in detect the disease and is thus insufficient to rule it out. In patients at increased risk, the sensitivity of the physical examination is even lower. However, it continues to be recommended once or twice a year for women aged 20–25 years of age and carrying pathogenic variants in *BRCA1, BRCA2, TP53* or *PTEN*.

For women in the breast cancer predisposition group, early mammographic screening is adopted, considering the earlier development of the disease, with the incorporation of complementary imaging tests such as MRI and ultrasound due to the limitations of the mammography examination for age groups below 40 years in the female population.

In the general population, mammography has shown to be related to a reduction in mortality rates, although its usefulness is less understood in women with pathogenic variants in *BRCA1, BRCA2, TP53, PTEN* and *STK11*, or with history of chest irradiation in the age of 10 to 30 years. In this group, annual mammography is recommended starting from 30 years old, with adjuvant MRI.

The guidelines related to the presence of pathogenic variants of moderate penetrance are less well-defined: annual mammography is recommended from the age of 40 onwards for patients with variants in *ATM, CHEK2* and *NBN*; and from the age of 30 onwards for cases of variants in *PALB2, CDH1* and *NF1*.

Although mammography remains an appropriate tool to screen the general population, its use alone may be insufficient to detect patients at increased risk of developing breast cancer. The method has less sensitivity in denser breasts, commonly present in younger patients, who constitute one of the groups considered at increased risk for hereditary breast cancer.

The MRI has a higher sensitivity compared to mammography to diagnose breast cancer in patients with hereditary predisposition. It is recommended annually from 25 to 30 years of age onwards in this group of women, also being considered annually from 30 to 50 years of age onwards — the age group in which mammography becomes the primary screening method.

A comparative analysis using a simulation model of pathogenic variants in *BRCA1* and *BRCA2* demonstrated that annual MRI from 25 years old onwards, accompanied by alternating digital mammography from the age of 30 onwards, is probably the most effective screening strategy, being related to the highest life expectancy.

According to data in the literature, when MRI and mammography were combined, the sensitivity goes up to 93%. Women with previous breast cancer are at greater risk of developing secondary tumors in the treated and contralateral breast; therefore, the combined use of imaging tests is also recommended.
Guidelines for MRI screening in women with moderate penetrance pathogenic variants are also not so well defined. Annual MRI is considered for patients with variants in ATM, CHEK2 and NBN from the age of 40 onwards, and the age of 30 onwards for patients presenting variants in PALB2, CDH1 and NF1.29

Recommendations for discontinuing MRI screening in patients at increased risk vary between age groups over 50 years old — except for patients with dense breasts —, and after the age of 75 or when the life expectancy of the patient is set at less than 10 years.10,29

Ultrasoundography, although not used as a routine method, can be useful as a complementary method in selected patients. Sensitivity is lower than that of MRI but comparable to that of mammography in young patients at increased risk. Therefore, in this group, it may be indicated mainly in women with dense breasts, pregnant women, lactating women or women who cannot undergo MRI.10

Although it does not provide many additional benefits in detecting cancer, the ultrasound can be used to increase the specificity of MRI by ruling out benign lesions. Furthermore, its adjuvant use may be more convenient and economical for short-term follow-up and also in guided biopsies.10

Although breast cancer is more common among women, men who carry mutations in the BRCA2 gene may be at increased risk of developing the disease. In this case, annual clinical breast exam and monthly self-examination are recommended from the age of 35 onwards. Due to the low incidence of breast cancer in this group, even in those at increased risk, there are no studies to determine the value of additional screening methods.29

### Genetic counseling and molecular research

Genetic counseling is a multifaceted process that can help to identify patients and family members who carry a mutation associated with increased risk of cancer. Genetic research should always be accompanied by pre- and post-test counseling, as to clarify all the possibilities of results, the limitations of the tests to be performed, and the possibilities of prevention, as well as to present the follow-up strategies and evaluate the chances of disease occurrence or recurrence in patients or relatives.34

International cancer research bodies propose guidelines that alert experts to pay attention to individuals at increased risk of hereditary cancers.10

According to guidelines by the National Comprehensive Cancer Network (NCCN), individuals who meet at least one of the following criteria should be referred for genetic counseling: personal history of breast and/or ovarian cancer; diagnosis under 50 years of age (in case of triple-negative breast cancer, personal history of two breast cancer diagnoses regardless of age of onset, and known mutation in a cancer-susceptibility gene within the family); several close family members with related cancers (breast, ovary, colon, endometrial, prostate, or pancreatic); diagnosis of breast cancer in men; and people of Ashkenazi Jewish ancestry with personal history of breast, ovarian and/or pancreatic cancer.10

In recent years, genetic testing has been allied to clinical practice. Until recently, the test was mainly performed by patients with a prominent family history of cancer encompassing a limited number of genes associated with a high or moderate risk of hereditary cancer. With the advent of the Next Generation Sequencing (NGS) molecular technique, panel genetic testing has become more widely used.36

Thus, there is scientific evidence of a clear association between hereditary cancer and some gene groups of high and moderate penetrance, with the presence of pathogenic variants that bring some possibilities of interference in therapeutics and disease management. In addition, the tracking of family members not yet affected by the disease is possible.10

The largest Brazilian study carried out by Palmero et al. sought to identify recurrent mutations in BRCA1 and BRCA2 that could be included in a low-cost genetic panel used as screening method for patients with predisposition to hereditary cancer. The study was carried out based on 649 genetic tests with pathogenic or probable pathogenic variants, obtained from 28 public and private health centers from 11 Brazilian states. In total, 126 mutations were identified in the BRCA1 gene and 103 in the BRCA2 gene, with 26 new variants identified in both genes.19

Table 1 lists some of the most prevalent mutations identified by the study.

However, some mutations were reported exclusively in certain geographic regions of the country, which suggests their founder effect and highlights the huge molecular heterogeneity and limited knowledge about these genes in the Brazilian population.19

### Table 1. Mutations identified by Palmero et al. in at least three probands

<table>
<thead>
<tr>
<th>BRCA1</th>
<th>n (%)</th>
<th>BRCA2</th>
<th>n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>c.5266dupC</td>
<td>89 (20.2)</td>
<td>c.2808_2811delACAA</td>
<td>20 (9.6)</td>
</tr>
<tr>
<td>c.3331_3334delCAAG</td>
<td>45 (10.2)</td>
<td>c.5946delT</td>
<td>15 (7.2)</td>
</tr>
<tr>
<td>c.68_69delAG</td>
<td>19 (4.3)</td>
<td>c.156_157insAlu</td>
<td>11 (5.3)</td>
</tr>
<tr>
<td>c.211A&gt;G</td>
<td>17 (3.9)</td>
<td>c.6405_6409delCTTAA</td>
<td>10 (4.8)</td>
</tr>
<tr>
<td>c.5074+2T&gt;C</td>
<td>14 (3.2)</td>
<td>c.2T&gt;G</td>
<td>8 (3.8)</td>
</tr>
<tr>
<td>c.470_471delCT</td>
<td>11 (2.5)</td>
<td>c.1138delA</td>
<td>7 (3.4)</td>
</tr>
<tr>
<td>c.1687C&gt;T</td>
<td>10 (2.3)</td>
<td>c.9382C&gt;T</td>
<td>7 (3.4)</td>
</tr>
<tr>
<td>c.4675+1G&gt;A</td>
<td>9 (2.0)</td>
<td>c.2266C&gt;T</td>
<td>3 (1.4)</td>
</tr>
</tbody>
</table>

Source: prepared by the authors based on data taken from the article by Palmero et al.19
To date, more than 35 candidate genes related to high and moderate risk of breast cancer have been suggested\(^9,28\). However, only few of these known genes had their variants significantly associated with breast cancer susceptibility, even in cases of positive family history or early diagnosis of the disease\(^9,37\).

By the same token, the use of multigene panels in genetic tests has considerably increased the number of patients diagnosed with a variant of uncertain significance (VUS), which reinforces the need for better models predictive of pathogenicity and increased efforts to help classify these variants, such as co-segregation analyses, personal and family history, co-occurrence of pathogenic variants, and histological and molecular characteristics of tumors\(^9,38\).

In order to better define the set of genes associated with breast cancer risk, Dorling et al. created a panel with 34 known genes that show an association with or susceptibility to breast cancer. The study included women with (60,466) and without breast cancer (53,461) from 25 countries who took part in population-based studies and studies based on families with a history of breast cancer, making up the Breast Cancer Association Consortium (BCAC)\(^36\).

Variants that cause alteration in protein function were associated with a significant risk of breast cancer (p<0.0001) in 5 genes: ATM, BRCA1, BRCA2, CHEK2 and PALB2 (95%CI 2.10–10.57). Susceptibility to breast cancer was also observed in 7 genes: BARD1 (OR=2.09; 95%CI 1.35–3.23), RAD51C (OR=1.93; 95%CI 1.20–3.11), RAD51D (OR=1.86; 95%CI 1.11–2.93), PTEN (OR=2.25; 95%CI 0.85–6.00), NF1 (OR=1.76; 95%CI 0.96–3.21), TP53 (OR=3.06; 95%CI 0.63–14.91) and MSH6 (OR=1.96; 95%CI 1.15–3.33) (Table 2)\(^36\).

Following similar objectives, the North American study conducted by Hu et al. involving 12 population-based studies used a panel with 28 breast cancer-predisposing genes evaluated in 32,247 case-patients and 32,544 control-patients (Table 3)\(^37,39\).

Pathogenic variants were identified in 12 genes established as predisposing to breast cancer in 5.03% of cases and 1.63% of controls. Corroborating the study by Dorling et al., the BRCA1 (OR=7.62; 95%CI 5.33–11.27) and BRCA2 (OR=5.23; 95%CI 4.09–6.77) genes are linked to a high risk for breast cancer; and the PALB2 (OR=3.83; 95%CI 2.68–5.63) and CHEK2 (OR=2.47; 95%CI 2.02–3.05) genes, to moderate risk\(^36\).

In women affected by the disease, the most prevalent mutations were observed in BRCA1 (OR=7.62; 95%CI 5.33–11.27), BRCA2 (OR=5.23; 95%CI 4.09–6.77) and PALB2 (OR=3.83; 95%CI 2.68–5.63). In unaffected women, most mutations were observed in CHEK2 and ATM, indicating a moderate risk for breast cancer\(^36\).

In summary, both studies showed a significant association between breast cancer risk and variants of 8 genes — BRCA1, BRCA2, PALB2, BARD1, RAD51C, ATM and CHEK2. However, most genes tested were not significantly associated with breast cancer, and the larger the multigene panel, the higher the VUS rates\(^36,39\).

Women who carry mutations in CHEK2 and ATM have tumors that express estrogen receptors, which may benefit from anti-estrogen therapies such as tamoxifen, raloxifene or aromatase inhibitors. However, studies involving chemoprevention have not been carried out in women with mutations in CHEK2 or ATM; and even among carriers of mutations in BRCA1 and BRCA2, the absorption of tamoxifen is low\(^40\).

With regard to other types of cancer, carriers of CHEK2 mutations are considered to be at high risk for colon cancer, and carriers of ATM mutations are considered at risk for pancreatic cancer.

### Table 2. The 34 genes in the study by Dorling et al.\(^36\)

| ABRAXAS1 | MSH2 |
| AKT1 | MSH6 |
| ATM | MUTYH |
| BABAM2 | NBN |
| BARD1 | NF1 |
| BRCA1 | PALB2 |
| BRCA2 | PIK3CA |
| BRIP1 | PMS2 |
| CDH1 | PTEN |
| CHEK2 | RAD50 |
| EPCAM | RAD51C |
| FANCC | RAD51D |
| FANCM | RECQL |
| GEN1 | RINT1 |
| MEN1 | STK11 |
| MLH1 | TPS3 |
| MRE11 | XRCC2 |

Source: prepared by the authors based on data taken from the article by Dorling et al.\(^19\)

### Table 3. The 28 genes in the study by Hu et al.\(^39\)

| ATM | MRE11A |
| BARD1 | MSH2 |
| BLM | MSH6 |
| BRCA1 | NBN |
| BRCA2 | NF1 |
| BRIP1 | PALB2 |
| CDH1 | PTEN |
| CDKN2A | RAD50 |
| CHEK2 | RAD51C |
| ERCC3 | RAD51D |
| FANCC | RECQL |
| FANCM | RINT1 |
| MLH1 | SLX4 |
| MRE11 | XRCC2 |
| TPS3 |

Source: prepared by the authors based on data taken from the article by Hu et al.\(^39\)
cancer. However, colon cancer screening is recommended for carriers of CHEK2 mutations, but not for ATM mutation carriers when it comes to pancreatic cancer in 10.

It is important to emphasize that the use of genetic panels and analyses of genomic rearrangements are great allies in the investigation of hereditarily-predisposed cancer, and that panels with multiple investigated genes must be well evaluated, as they can generate data for which clinical management has not yet been determined and, therefore, some patients may choose to have a genetic test with a smaller panel of genes, containing only high and moderate risk genes, as these provide the best-characterized cancer risk estimates and management recommendations41.

Table 4 presents nine genes associated with breast cancer risk, including estimated lifetime risk, other malignancies associated with the presence of gene mutations, and individualized management and screening approaches according to NCCN guidelines28, as well as data presented in studies by Dorling et al.36, Bharucha et al.42, Owens et al.35 and Shiovitz et al.28.

<table>
<thead>
<tr>
<th>Germline mutation</th>
<th>Cumulative risk* of breast cancer</th>
<th>Other associated malignancies</th>
<th>Screening guidelines†</th>
</tr>
</thead>
<tbody>
<tr>
<td>BRCA1</td>
<td>85%</td>
<td>Ovary, fallopine tubes, peritoneum, pancreas, prostate, colon.</td>
<td>25–29 years*: Annual MRI (mammography if MRI is not available); 30–75 years: annual mammography and MRI; discuss preventive mastectomy and bilateral salpingo-oophorectomy after pregnancy.</td>
</tr>
<tr>
<td>BRCA2</td>
<td>65%</td>
<td>Ovary, fallopine tubes, peritoneum, pancreas, prostate, melanoma.</td>
<td>25–29 years*: Annual MRI (mammography if MRI is not available); 30–75 years: annual mammography and MRI; discuss preventive mastectomy and bilateral salpingo-oophorectomy after pregnancy.</td>
</tr>
<tr>
<td>TP53</td>
<td>85%</td>
<td>Sarcomas and CNS, adrenocortical, gastrointestinal, and associated with radiation.</td>
<td>25–29 years*: Annual MRI (mammography if MRI is not available); 30–75 years: annual mammography and MRI; discuss bilateral preventive mastectomy.</td>
</tr>
<tr>
<td>PTEN</td>
<td>67–87%</td>
<td>Thyroid, endometrium, colorectal, renal.</td>
<td>30–35 years*: Annual MRI and mammography; discuss preventive mastectomy.</td>
</tr>
<tr>
<td>CDH1</td>
<td>42–60%</td>
<td>Diffuse gastric cancer.</td>
<td>30 years or older: annual mammography, consider annual MRI; insufficient evidence for preventive mastectomy.</td>
</tr>
<tr>
<td>STK11§</td>
<td>44–50%</td>
<td>Colorectal, stomach, small intestine, pancreas, ovary, Sertoli cell tumor.</td>
<td>25–29 years*: Annual MRI; 30 years and older: annual mammography and MRI.</td>
</tr>
<tr>
<td>ATM</td>
<td>20%</td>
<td>Pancreas.</td>
<td>≥40 years: annual mammography; consider annual MRI; insufficient evidence for preventive mastectomy, or prescription of radiation therapy.</td>
</tr>
<tr>
<td>CHEK2</td>
<td>20–25%</td>
<td>Colorectal, stomach, prostate, kidney and thyroid.</td>
<td>≥40 years: annual mammography; consider annual MRI; insufficient evidence for preventive mastectomy.</td>
</tr>
<tr>
<td>PALB2</td>
<td>33%–59%</td>
<td>Pancreas.</td>
<td>≥30 years: annual mammography and MRI; insufficient evidence for preventive mastectomy.</td>
</tr>
</tbody>
</table>

MRI: magnetic resonance imaging; CNS: central nervous system. *Estimated cumulative risk up to 70 years of age; †Due to a lack of local studies, the recommendations in Brazil are based on international data; ‡Or 10 years before the youngest person affected in the family; §Gene evaluated in the study by Dorling et al.36, but without statistical significance of risk. Source: Prepared by the authors, based on information taken and adapted from10,28,35,36,42.
Therapeutic and risk-reducing approaches in hereditary breast cancer

The term “risk reduction” has been considered more appropriate than “prophylactic” in recent times, as no mastectomy can remove all of the breast tissue. Studies have shown a reduction in breast cancer risk of approximately 95% in BRCA1 and BRCA2 mutation carriers undergoing bilateral risk-reducing mastectomy (BRRM) in combination with oophorectomy, and a reduction in risk of approximately 90% in those with intact ovaries46-48.

A recent systematic review confirms the benefit of BRRM in reducing breast cancer incidence and mortality in patients at high risk for breast cancer predisposition such as carriers of BRCA1 and BRCA2 mutations, but this evidence requires rigorous prospective studies due to methodological flaws in the existing literature49. Contralateral mastectomy as risk reduction (CMRR) data for patients who had unilateral breast cancer are not conclusive, as existing studies show a reduction in the incidence of contralateral breast cancer but no definitive survival benefit50-53.

The main treatment strategies for breast cancer are surgery and systemic treatment. One of the main concerns in the surgical treatment of breast cancer with a pathogenic variant for BRCA1 and BRCA2 is whether the outcome of treatment with breast-conserving surgery (BCS) combined with adjuvant radiotherapy is equivalent to radical mastectomy54.

A study compared results of the surgical method (BCS combined with radiotherapy versus mastectomy) in cases of breast cancer with pathogenic variants BRCA1 and BRCA2. According to the work by Onitilo et al., a higher 10-year survival was observed in the group undergoing BCS with adjuvant radiotherapy (BCS: 80.9% versus mastectomy: 67.2%), in addition to lower rates of local recurrence55.

However, it is known that women who carry mutations in BRCA1 and BRCA2 are more likely to develop a secondary cancer, that is, ipsilateral or in the contralateral breast. For these patients, a bilateral mastectomy is recommended, as studies suggest that women who carry mutations in BRCA1 and BRCA2 and who undergo bilateral mastectomy are less likely to die of breast cancer than women who have been treated with unilateral mastectomy56-52.

A meta-analysis encompassing 526 patients with a pathogenic variant in BRCA1 and BRCA2 and 2,320 patients with sporadic breast cancer showed no difference in overall survival rates between these groups. However, patients with mutations in BRCA1 and BRCA2 had a greater recurrence of ipsilateral breast cancer than patients with sporadic breast cancer, with a mean follow-up of more than six years (RR=1.51; 95%CI 1.15–1.98)52.

Radiation after BCS is not performed only in very exceptional cases. Given the essential role of the BRCA1 and BRCA2 genes in DNA repair of other cancer-inducing genes in humans, questions have been raised regarding the possible complications of radiotherapy in breast cancer involving pathogenic variants in BRCA1 and BRCA255. However, a study by Pierce et al. showed no significant difference in radiation complication rates between women carrying BRCA1 and BRCA2 mutations versus women with sporadic cancer56.

In this setting, radiotherapy also plays an important role after mastectomy. Indications should be similar in both radical surgery and conservative mastectomy. Traditionally, radiotherapy is indicated for patients with four or more affected lymph nodes, positive surgical margins, or with tumors larger than 5 cm. However, there is a debate about the role of radiotherapy in patients with 1–3 metastatic lymph nodes and the role of secondary factors such as age, molecular subtype and angiolymphatic invasion in the decision-making about the use of radiotherapy after mastectomy, remaining quite controversial58.

In patients with the Li-Fraumeni syndrome presenting with germline mutations in the TP53 gene, exposure to radiotherapy increases the risk of a second cancer. In these patients there is an inactivation of DNA repair mechanisms and activation of apoptosis, so the susceptibility to radio-induced tumors can accelerate the appearance of a second neoplasm59.

The repair pathway by homologous recombination of damaged DNA — in which there is loss of function caused by mutations present in the BRCA1 and BRCA2 genes — lead to very similar phenotypes, which fall within the hereditary predisposition to breast and ovarian cancer. Likewise, mutations in RAD51C, BRIP1, PALB2 and others can lead to a phenotype similar to that of HBOC58.

By taking into account the chemotherapy-based treatment, which causes DNA damage requiring repair genes of the homologous recombination pathway to induce a repair response, the status of the pathogenic variant in BRCA1 and BRCA2 is considered a decisive factor to predict sensitivity to chemotherapy60.

The profile of genomic structural alterations caused specifically by homologous recombination deficiency (HRD) repair has been studied as potential markers of pathway deficiency through scores, which may be useful in evaluating the association not only with the response rate to chemotherapy, but also with clinicopathological and overall survival factors59.

In vitro studies, cells with the BRCA1 variant were shown to be more sensitive to platinum-based chemotherapeutic agents, as they disrupt the DNA structure. They also showed greater resistance to microtubule-inhibiting chemotherapies such as taxanes. These findings were supported by data from patients with BRCA1 and BRCA2 pathogenic variant breast cancer who underwent palliative or neoadjuvant taxane-only chemotherapy61. However, there is insufficient evidence to exclude taxanes from adjuvant chemotherapy strategies in patients with breast cancer carrying mutations in BRCA1 and BRCA262.

Patients diagnosed with breast cancer at younger ages and carriers of mutations in high and moderate penetrance genes should have an individualized surgical treatment. Carriers of
BRCA1 and BRCA2 mutations, for example, face more aggressive surgical interventions for therapeutic purposes and to reduce the risk of developing primary or contralateral breast cancer, which is increased57. However, breast-conserving surgery, as well as skin-sparing mastectomies with or without preservation of the nipple-areolar complex, have been shown safe and to provide a better restoration. Selecting the best surgical approach for this group of patients requires taking into account several factors, including genetic risk, personal and family history, and the patient’s own preferences57.

AUTHORS’ CONTRIBUTIONS

REFERENCES


