Hereditary breast and ovarian cancer susceptibility genes (Review)

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Abstract. Women with hereditary breast and ovarian cancer (HBOC) syndrome represent a unique group who are diagnosed at a younger age and result in an increased lifetime risk for developing breast, ovarian and other cancers. This review integrates recent progress and insights into the molecular basis that underlie the HBOC syndrome. A review of English language literature was performed by searching MEDLINE published between January 1994 and October 2012. Mutations and common sequence variants in the BRCA1 and BRCA2 (BRCA) genes are responsible for the majority of HBOC syndrome. Lifetime cancer risks in BRCA mutation carriers are 60-80% for breast cancer and 20-40% for ovarian cancer. Mutations in BRCA genes cannot account for all cases of HBOC, indicating that the remaining cases can be attributed to the involvement of constitutive epimutations or other cancer susceptibility genes, which include Fanconi anemia (FA) cluster (FANCD2, FANCA and FANCC), mismatch repair (MMR) cluster (MLH1, MSH2, PMS1, PMS2 and MSH6), DNA repair cluster (ATM, ATR and CHK1/2), and tumor suppressor cluster (TP53, SKT11 and PTEN). Sporadic breast cancers with TP53 mutations or epigenetic silencing, ER- and PgR-negative status, an earlier age of onset and high tumor grade resemble phenotypically BRCA1 mutated cancers termed 'BRCAness', those with no BRCA mutations but with a dysfunction of the DNA repair system. In conclusion, genetic or epigenetic loss-of-function mutations of genes that are known to be involved in the repair of DNA damage may lead to increased risk of developing a broad spectrum of breast and ovarian cancers.

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1. Introduction

Breast cancer and ovarian cancer develop through multiple molecular pathways guided by genetic and epigenetic clonal selections. Both cancers have been recognized as a heterogeneous disease with regard to clinical and biological properties. The majority of cancer cases are considered sporadic-appearing tumors in nature because there is no obvious family history, but a number of families with a known genetic cause or inherited predisposition to cancer have been identified (1). Individuals who carry an inherited genetic mutation and epigenetic aberrations in the tumor suppressor genes have an increased lifetime risk of developing cancer. Germline mutations in cancer susceptibility genes cause cancer if the wild-type allele is lost or inactivated. Breast and ovarian cancers (5-10%) may be hereditary and occur in cancer prone syndromes (2-7). Patients predisposed to breast and ovarian cancers are known as the hereditary breast and ovarian cancer (HBOC) syndrome.

HBOC syndrome is an autosomal dominantly inherited disease characterized by a young age of onset, more than one synchronous or metachronous tumor, and a family history of first and second degree relatives with similar cancers (8). Mainly HBOC syndrome results from germline mutations in breast cancer genes BRCA1 or BRCA2. Other genes or low penetrance alleles might be associated with the HBOC phenotype (9). There is an increasing understanding that the interrelationship between BRCA gene cluster and Fanconi anemia (FA), mismatch repair (MMR) and DNA repair gene status plays a key role in the pathogenesis of cancer predisposition syndromes. We reviewed the HBOC syndrome and current knowledge of inherited susceptibility genes.

2. Materials and methods

A computerized literature search was performed to identify relevant studies reported in the English language. We searched MEDLINE electronic databases (http://www.ncbi.nlm.nih.gov/sites/entrez) published between January 1994 and
October 2012, combining the keywords ‘hereditary breast and ovarian cancer’, ‘pathogenesis’, ‘BRCA’ and ‘DNA repair’. Various combinations of the terms were used, depending on the database searched. Each gene was also linked to NCBI Entrez Gene pages (http://www.ncbi.nlm.nih.gov/sites/entrez). Additionally, references in each article were searched to identify potentially missed studies.

3. Current understanding of HBOC susceptibility genes

**BRCA1 and BRCA2**. The breast cancer-associated genes BRCA1 on chromosome 17q and BRCA2 on chromosome 13q are the most well-known breast cancer susceptibility genes (4,5). The HBOC syndrome is linked to the BRCA1 gene and, to a lesser extent, to the BRCA2 gene. Germline mutations in these genes account for 2.5% (up to 10%) of all breast cancers and all ovarian cancers. Mutations were present with a frequency of more than 10% in the high risk populations, including patients with a family history of breast or first-degree ovarian cancer, those with bilateral breast cancer, multiple organ cancer including younger breast cancer patients (aged <35 years). Mutations in the BRCA1 and BRCA2 genes are responsible for ~60% (up to 85%) of HBOC. Women carrying a BRCA1 or BRCA2 genetic mutation have 60-80% and 20-40% lifetime risk of developing breast cancer and ovarian cancer, respectively. More modest increases in risk for other cancers have been noted: additional sites included stomach, pancreas, prostate and colon. The cancer risk ranged from 20 to 60%, with the greatest increases in cancer risk in stomach and pancreas. BRCA mutations were also associated with increased risks for leukemia and lymphoma (10).

A recent study showed that breast cancer patients belonging to a population with a high probability of being BRCA1 carriers showed a better prognosis compared with those with sporadic breast cancer (11). Furthermore, BRCA1 and BRCA2-related invasive epithelial ovarian cancers have a better 5-year overall survival compared with sporadic ovarian cancers (12). The 5-year overall survival was better in BRCA2 carriers compared to BRCA1 carriers (36% for non-carriers, 44% for BRCA1 carrier and 52% for BRCA2 carriers).

The mutations include partial or complete gene deletions, duplications, large insertions, splice alteration, frameshifts as well as missense and nonsense mutations. Deletions or insertions usually lead to abnormal structure and function. Germline mutations are usually pathogenic point mutations, and are scattered throughout their coding regions. The potential hot-spot mutations within BRCA 1 and BRCA2 are uncommon. The previously described mutations were identified in the Breast Cancer Information Core website (BIC, http://research.nhghri.nih.gov/bic/). More than 3500 mutations have been reported throughout both genes. Data from subjects with a variety of ethnic backgrounds altered the overall odds of BRCA mutation carrier status. The spectrum of mutations is different depending on the race. The mutations are detected in 10-12% of Ashkenazi Jewish women diagnosed with breast cancer. Ashkenazi Jewish subjects are observed at increased frequency compared to other Caucasian, because this population harbors ancient BRCA1 and BRCA2 mutant alleles. Mutation analysis revealed the c. 5266dup (until recently referred in the literature as 5382insC), c. 68_69del (185delAG) and 4153delA mutations in BRCA1 and c. 5946del (6174delT) mutation in BRCA2 (13). BRCA gene founder mutations such as BRCA1c. 5266dup mutation, BRCA2999del5 mutation and BRCA1delexon17 have also been described in other populations, including the Slavic, Finnish, Icelandic and German populations, respectively.

The tumor suppressor BRCA1 and BRCA2 genes are essential components of the double-strand break (DSB) repair by homologous recombination (HR) system. Targeting tumor suppressor loss-of-function is possible based on the concept of synthetic lethality. Thus, the synthetically lethal effect might be observed in tumors defective in BRCA1 or BRCA2 that are required for efficient HR, indicating that ovarian cancer patients carrying germ-line mutations had improved rates of progression-free and overall survival (12). Chromosomal rearrangements might be formed as a consequence of these error-prone DSB repairs and lead to the development of genomic instability. Large genomic rearrangements have been identified in HBOC families and account for 8-15% of deleterious BRCA mutations, but these rearrangements may escape detection (14). The issue is that BRCA genetic testing done through sequencing will not capture large rearrangements in these genes.

BRCA1 functions as a tumor suppressor gene, but paradoxically, BRCA1 knockout mice are embryonically lethal in homozygous state. Lack of BRCA1 is thought to result in cellular lethality, suggesting that BRCA1 regulates stem/progenitor cell proliferation and differentiation. For cell differentiation, BRCA1 regulates apicobasal polarity, together with several genes such as RHAMM (hyaluronan-mediated motility receptor), AURKA (aurora kinase A) and TPX2 (microtubule-associated, homolog). Intracellular RHAMM associates with BRCA1 and BARD1 (BRCA1 associated RING domain 1). The complex attenuates the mitotic-spindle-promoting activity of RHAMM that might contribute to tumor progression. BRCA1 also binds and regulates AURKA, a cell cycle-regulated kinase that appears to be strongly involved in centrosome regulation. Genetic variants in the AURKA gene may contribute to breast cancer development (5). BRCA1 further accumulates TPX2 and is required for mitotic spindle-pole assembly. BRCA-associated nuclear core complex proteins are required for the functional integrity of the pathway of not only DNA damage response and repair, but also cell differentiation.

From a clinical point of view, the triple-negative breast cancer characterized by the absence of estrogen receptor (ER), progesterone receptor (PgR), and HER2 (also known as ERBB2) accounts for ~15% of breast cancers and is diagnosed more frequently in younger women (15). Compared with the ER/PR-positive tumors, triple-negative breast cancer showed a greater risk for recurrence and shortened survival. Triple-negative breast cancer was frequently associated with mutations in BRCA genes: the incidence was 12.5-20% (16). Women with an early age-of-onset triple-negative breast cancer are more likely to be associated with deleterious mutations in BRCA1 and BRCA2 genes (17). Even in non-BRCA gene mutations, a subset of triple-negative tumors shares multiple clinicopathologic features and phenotype with BRCA-mutated breast cancers. They harbor dysfunctional DNA repair mechanisms, but the nature of this link remains opaque. A subset
of HBOC syndrome also contain mutations in the TP53 gene, and the TP53 loss-of-function tumors have a low frequency of HER2 expression (18).

The predominant histologic type of ovarian cancers associated with the HBOC syndrome was high-grade serous carcinomas of the ovary. There were no significant differences in ovarian cancer morphology between BRCA1 and BRCA2 carriers (19). Ovarian cancer patients with BRCA mutations were associated with an increased chemosensitivity and improved overall survival, but some investigators failed to confirm improved survival.

Modifying hereditary breast and ovarian cancer risks. Mutations in BRCA1 and BRCA2 do not account for all cases of HBOC, implicating that the remaining cases can be attributed to the involvement of other susceptibility genes. Other genes, including Fanconi anemia (FA) cluster (FANCD2, FANCA and FANCN), MMR cluster (MLH1, MSH2, PMS1, PMS2 and MSH6), DNA checkpoint cluster (ATM, ATR and CHK1/2), and tumor suppressor cluster (TP53, SKT11 and PTEN) have been associated with increased risk of breast and ovarian cancer as part of other cancer syndromes. The contribution of mutations in other genes to the burden of breast or ovarian cancer is indicated in Table I.

Poly(ADP-ribose) polymerase (PARP). Poly(ADP-ribose) polymerase (PARP) is an enzyme involved in the recovery of cells from DNA damage and the regulation of the molecular events such as BER, a key pathway in the repair of DNA single-strand breaks (SSB). The inhibition of PARP leads to the induction of synthetic lethality and cell death by targeting HR-mediated DNA repair deficient tumors (20). Tumors that lack functional BRCA1, BRCA2, or TP53 are hypersensitive to inhibition of PARP. Several proteins involved in HR on sensitivity to PARP inhibition may include BRCA cluster (RAD51C, RAD51D and RAD54), FA cluster (FANCD2, FANCA and FANCN), Cdk cluster, nucleotide excision repair (NER) cluster (RPA1 and NBN), DNA repair checkpoint cluster (ATR, ATM, CHK1 and CHK2) and TP53 cluster. Therefore, therapeutic approach using PARP inhibitors may be feasible for BRCA dysregulated tumors and appear promising in a variety of cancer types, including breast and ovarian cancers. The presence of these germline mutations and epimutations types might be a hallmark of BRCAness and a potential biomarker for sensitivity to PARP inhibition.

Recently, Kaye et al (21) reported the results of a phase II study comparing the efficacy and safety of olaparib, a potent oral PARP inhibitor, in patients with germ-line BRCA mutations and recurrent ovarian cancer. Unfortunately, there was no remarkable differences in PFS between the olaparib and pegylated liposomal doxorubicin groups. Another clinical study demonstrated that olaparib applied as maintenance treatment prolonged PFS, but not OS, in patients with advanced and platinum-sensitive, recurrent, high-grade serous ovarian cancer (22).

BER and NER. There are two types of DNA repair proteins: the nucleotide excision repair (NER) pathway (ERCC1, excision repair cross-complementing rodent repair deficiency, complementation group 1) and the base excision repair (BER) pathway (XRCC1, X-ray repair complementing defective repair in Chinese hamster cells 1) (23). The NER pathway is responsible for the removal of bulky DNA lesions. NER is a defense system against various types of DNA damage and necessary for maintaining genomic stability. RPA1, replication protein A1, is a single-stranded DNA binding protein and participates in the recruitment of the two structure-specific DNA endonucleases, XPG (xeroderma pigmentosum, also known as ERCC5) and XPF (xeroderma pigmentosum, complementation group F)-ERCC1 complex, which makes the 5' incision in NER. The XPF-ERCC1 complex is essential for cutting the damaged DNA strand and the DNA repair by the NER pathway. NBN (nibrin, also known as NBS1) was involved in DNA DSB repair and DNA damage-induced checkpoint activation as a component of the MRE11-RAD50-NBS1 complex. Mutations in NBN is thought to be associated with breast-cancer risk.

Furthermore, XRCC1 is involved in the repair of DNA SSB and oxidative damage (23). This protein interacts with DNA ligase III, polymerase β and PARP1 participating in the BER pathway. Reduced XRCC1 expression may confer chemoradiation and leads to improved patient survival.

4. Epigenetic silencing

Between 50-80% of HBOC syndrome can be explained by defective germline mutations in BRCA1 and BRCA2 as well as, to a lesser degree, other genes described above, but for the remaining families the factors driving susceptibility remain unknown (24). Approximately one third of the HBOC families do not have evidence of the germline mutations in BRCA1 and BRCA2. The loss of BRCA function might be due to either germline/somatic mutation or epigenetic silencing. Since little is known about the contribution of epimutations to the remaining BRCA1/2 mutation-negative cases, epigenetic silencing has been explored in HBOC syndrome. The activities of tumor suppressor genes and cancer susceptibility genes could be influenced by genetic and epigenetic alterations. Decreased expression of cancer susceptibility genes has been observed in sporadic breast and ovarian cancer where it is often associated with the aberrant epimutations or hypermethylation of the BRCA1 and BRCA2 genes. The loss of BRCA1 function due to somatic hypermethylation explained ~10% of sporadic breast cancer cases. A subset of the sporadic tumor patients demonstrated hypermethylation of BRCA2 and their interacting protein including HP1γ (heterochromatin protein 1gamma), RAD51C, ATM and PALB2 (25). Based on BRCA1 deletion, TP53 mutations, ER- and PgR-negative status, young age at diagnosis and high grade tumor, phenotypic features of sporadic breast cancers resemble BRCA1 mutated cancers termed ‘BRCAness’. Phenotypic similarities were most closely observed in BRCAness, epigenetic silencing and deletion of the BRCA1 and BRCA2 genes.

5. Lynch syndrome

Many genes have been implicated in the DNA damage response pathways where the BRCA1 and BRCA2 genes are involved. Genetic susceptibility to cancer is attributed to deleterious germline mutations in the DNA mismatch repair (MMR) genes. Another important non-BRCA1/BRCA2
### Table I. Modifying hereditary breast and ovarian cancer risks.

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<th>No.</th>
<th>Genes</th>
<th>Functions</th>
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<td><strong>Genes with high penetrance</strong></td>
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<tr>
<td>1</td>
<td>CDH1, cadherin 1 type 1</td>
<td>Loss of CDH1 function contributes to progression in cancer by increasing proliferation, invasion, metastasis and epithelial-mesenchymal transition (EMT). Germline mutations in the CDH1/E-cadherin gene predispose to the development of the autosomal hereditary diffuse gastric cancer (HDGC) syndrome, which leads to the development of breast, colorectal, thyroid and ovarian cancer (46). Gastric cancer risk is also elevated in HBOC, in Lynch syndrome (due to germline mutations in DNA mismatch repair genes), in familial adenomatous polyposis (FAP) [germline adenomatosis polyposis coli (APC) mutations], in Li-Fraumeni syndrome (germline p53 mutations), in Peutz-Jeghers syndrome (germline STK11 mutations), and in juvenile polyposis syndrome [germline mutations in the SMAD family member 4 (SMAD4) and bone morphogenetic protein receptor, type IA genes (BMPR1A)] (47).</td>
<td>(46,47)</td>
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<td>2</td>
<td>NBS1, also known as NBN (nibrin)</td>
<td>The encoded protein nibrin is an integral member of the MRE11/RAD50/NBN (MRN) complex essential for processing DNA double-strand breaks and DNA damage-induced checkpoint activation. Mutations in this gene causes Nijmegen breakage syndrome, an autosomal recessive chromosomal instability syndrome characterized by microcephaly, growth retardation, immunodeficiency, and cancer predisposition. The NBN 657del5 mutation plays a role in breast cancer risk, but not ovarian cancer risk.</td>
<td>(47)</td>
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<td>3</td>
<td>NF1, neurofibromin 1</td>
<td>NF1 functions as a negative regulator of the Ras/Raf/Erk signal transduction pathway. Mutations in this gene are associated with neurofibromatosis type 1 (NF1) and other neoplasms, including juvenile myelomonocytic leukemia (JMML) and breast cancer (48).</td>
<td>(48)</td>
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<td>4</td>
<td>PTEN, phosphatase and tensin homolog</td>
<td>PTEN, a tumor suppressor gene, is mutated in a large number of cancers at high frequency. PTEN is a phosphatidylinositol 3-phosphatase and functions as a tumor suppressor by negatively regulating AKT signaling pathway. Mutations in PTEN gene are associated with the development of Cowden syndrome and also correlated with hamartomatous polyps, early-onset breast, thyroid and endometrial cancers (49).</td>
<td>(49)</td>
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<td>5</td>
<td>TP53, tumor protein p53</td>
<td>The encoded protein p53 responds to diverse cellular stresses to maintain genetic stability, thereby inducing cell cycle arrest, apoptosis, senescence, DNA repair, or changes in metabolism. Mutations, loss of heterozygosity (LOH), overexpression and loss of TP53 function occur as somatic mutations in human malignancies at high frequency. TP53 loss-of-function is essential for BRCA1-associated tumorigenesis (18). The mutation frequency of p53 in the three groups were sporadic ovarian cancers, 50%; somatic BRCA1 mutations, 90% and germline BRCA1 mutations, 80% (50). Germline mutations in TP53 are associated with hereditary cancers such as Li-Fraumeni syndrome, characterized by predisposition to multiple cancers.</td>
<td>(18,50)</td>
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<td>6</td>
<td>STK11, serine/threonine kinase 11</td>
<td>STK11 regulates cell polarity and functions as a tumor suppressor. Germline mutations in STK11 gene is associated with Peutz-Jeghers syndrome, an autosomal dominant disorder characterized by the growth of polyps in the gastrointestinal tract, pigmented macules on the skin and mouth, and cancer susceptibility in various organs including testis, ovary, endocervix, breast, pancreas and colon.</td>
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<td><strong>Genes with moderate penetrance</strong></td>
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<td>7</td>
<td>ATM, ataxia telangiectasia mutated</td>
<td>The protein encoded by this gene belongs to the PI3/P4-kinase family. The activation of ATM was mediated via interactions with a wide variety of downstream proteins, including checkpoint kinase CHK2, checkpoint proteins RAD17 and RAD9, and DNA repair protein NBN (51). ATM-mediated CHK2 kinase phosphorylation activates a variety of downstream substrates, including the CDC25C (cell division cycle 25 homolog C), TP53, and BRCA1 gene products, which are required for the induction of G2/M cell cycle arrest.</td>
<td>(6,51,52)</td>
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<td>7</td>
<td>ATM, ataxia telangiectasia mutated</td>
<td>and apoptosis after DNA damage. Germline mutations in this gene are associated with ataxia telangiectasia (AT), an autosomal recessive disorder. Polymorphisms in the ATM gene had an evidence for an association between variants in candidate genes and risk of breast cancer (6). A heterozygous variant, IVS10-6T&gt;G, was reported in breast cancer families (52). This variant seems to occur at a lower frequency (0.83%) in affected individuals. In contrast to BRCA1 and BRCA2, the CHK2 heterozygosity does not increase the ovarian cancer risk.</td>
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<td>8</td>
<td>ATR, ataxia telangiectasia and Rad3 related</td>
<td>ATR also activates several important key proteins, including checkpoint kinase CHK1, TP53, BRCA1, RAD17 and RAD9. Constitutively phosphorylated CHK1 could have activated oncogene signaling pathways including myc signaling. CHK1 phosphorylates the FANCE subunit of the Fanconi anemia (FA) core complex, which is associated with the FA-BRCA pathway (53).</td>
<td>(53)</td>
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<td>9</td>
<td>CHEK2, checkpoint kinase 2</td>
<td>CHEK2, a cell cycle checkpoint regulator, is essential for activation in response to DNA damage. CHEK2 inhibits CDC25C phosphatase, stabilizes p53, and leads to cell cycle arrest in G1. Since mutations in CHEK2 gene such as c.470T&gt;C and c.1100delC increase the risk of HBOC, CHEK2 has been identified as an breast cancer susceptibility gene in the high-risk BRCA1/2-founder mutation-negative individuals (54).</td>
<td>(54)</td>
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<td>10</td>
<td>Fanconi anemia (FA) genes</td>
<td>Mutations of Fanconi anemia (FA) genes have been demonstrated to cause breast and ovarian cancer susceptibility in cases from non-BRCA1/BRCA2 families (55). FA is a rare autosomal or X-linked recessive genetic heterogeneous disorder characterized by spontaneous chromosomal breaks, abnormal DNA repair and clinical problems including congenital abnormalities, endocrinopathies, early onset bone marrow failure and an increased susceptibility to cancer and leukemia (36). Similar to BRCA, the FA proteins form a functional core complex and are associated with a novel pathway required for DNA damage repair, particularly DNA interstrand cross-links. Owing to the link between FA and BRCA genes, this pathway is often referred to as the FA-BRCA pathway. FA proteins interact with each repair machinery such as that involved in HR, base excision repair (BER), nucleotide excision repair (NER), or MMR complex. As DNA damage response, the pathway-specific driver proteins have been shown to co-localize in nuclear foci with the FA core complexes such as FANCJ (BACH1)-BRCA1 and FANCN (PALB2)-BRCA1/2.</td>
<td>(36,55)</td>
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<td>11</td>
<td>FANCJ, Fanconi anemia group J protein, also known as BRCA1 interacting protein, C-terminal helicase 1</td>
<td>FANCJ is the BRCA1 associated C-terminal helicase (BACH1). The BRCA1-FANCJ interaction is required for promoting error-free repair, DNA DSB repair, and interstrand cross-links repair by linking to MMR protein complex MLH1-PMS2 (MutL α) and checkpoint control. Tumors associated with FANCJ mutations will compromise error free NHEJ (non-homologous end-joining) and resemble BRCA1-associated cancer (56).</td>
<td>(56)</td>
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<td>12</td>
<td>PALB2, partner and localizer of BRCA2</td>
<td>PALB2 (the partner and localizer of BRCA2), also known as FANCN, was identified as a BRCA1- and BRCA2-interacting protein. PALB2 is associated with the formation of the BRCA1-PALB2-BRCA2 complex and has rare, but moderate-risk for breast cancer (57). FANCD1 was identical to BRCA2. The FA complex is also required for the monoubiquitination of FANCD2. Furthermore, FANCD2 interacts with the MMR complex proteins MSH2 and MLH1. FANCD2 inactivation confers a hypersensitivity towards DNA cross-links and oxidative stress (58). Increased sensitivity to DNA cross-linking agents was attributable to defective MSH2 or MLH1 function. Recent evidence</td>
<td>(57,58)</td>
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<td>12</td>
<td>PALB2, partner and localizer of BRCA2</td>
<td>indicated that mutations in FANCD1 (BRCA2), FANCD2, FANCJ and FANCP have a role in breast and ovarian cancer susceptibility. Although clinical manifestations vary widely from patient to patient, individuals with FA are predisposed to develop breast and ovarian cancer than those without FA.</td>
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<td>13</td>
<td>CDK1, cyclin-dependent kinase</td>
<td>Cyclin-dependent kinase (CDK) is the driver for cell cycle progression. CDK was generally suppressed in response to DNA damage, but certain CDK seems necessary for proper DNA damage response. CDK1 phosphorylates BRCA1, and is essential for efficient formation of BRCA1 foci (59). CDK1 governs the G2/M transition. Depletion or inhibition of CDK1 results in deficient DNA damage response signaling and DNA repair by HR, suggesting that CDK1 may interfere with DNA damage response. As expected, combined inhibition of CDK1 and PARP in BRCA-wild-type cancer cells resulted in efficient tumor regression in a preclinical model (59).</td>
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<td>14</td>
<td>RAD50, RAD50 homolog</td>
<td>Other predisposition genes involved in DNA repair may account for a proportion of HBOC families. RAD genes are involved in DNA DSB repair and required for both types of the repair processes, NHEJ and HR (60). Polymorphic variants and defective mutations of RAD50 may be at a risk of developing breast cancer possibly through the loss-of-function of the MRE11-RAD50-NBN complex, key factors in maintaining genome stability (61). Some patients with an HBOC-like phenotype have mutations in RAD50. Mutations in this gene are the cause of Nijmegen breakage syndrome-like disorder. Heterozygous germline mutations in RAD51C were found in breast and ovarian cancer families (62). Pathogenic RAD51C mutations have also been identified in BRCA1/2 mutation-negative HBOC families (63). A biallelic mutation or a homozygous missense mutation in RAD51C was reported to cause FA-like disorder.</td>
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<td>15</td>
<td>FGFR2, fibroblast growth factor receptor 2</td>
<td>Mutations in the FGFR2 gene cause several disorders, including Crouzon syndrome, Pfeiffer syndrome, craniosynostosis, Apert syndrome, Jackson-Weiss syndrome, Beare-Stevenson cutis gyrata syndrome, Saethre-Chotzen syndrome, and syndromic craniosynostosis. The rs2981582, rs2420946, and rs1219648 FGFR2 polymorphisms act as modifiers of breast cancer susceptibility, particularly in the group of non-carriers of BRCA1/2 mutations (64).</td>
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<td>16</td>
<td>LSP1, lymphocyte-specific protein 1</td>
<td>LSP1 encodes an intracellular F-actin binding protein and regulates LS neutrophil motility, adhesion to fibrinogen matrix proteins, and transendothelial migration. The LSP1 rs3817198 polymorphism is associated to a modified risk of breast cancer.</td>
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<td>17</td>
<td>MAP3K1, mitogen-activated protein kinase kinase 1 E3 ubiquitin protein ligase</td>
<td>MAP3K1 is associated with activation of the ERK and JNK kinase pathways as well as the NF-kB pathway. Functional polymorphism of MAP3K1 rs889312 has been shown to influence the risk of familial and early-onset breast cancer.</td>
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<td>18</td>
<td>TGFB1, transforming growth factor, β1</td>
<td>Mutations in the TGFB1 gene is a genetic risk factor for Camurati-Engelmann disease featuring histopathological changes of osteomalacia. The rs1982073 TGFB1 polymorphism has been implicated in an elevated risk of progesterone receptor negative breast cancer.</td>
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<td>19</td>
<td>TOX3, TOX high mobility group box family member 3</td>
<td>TOX3 is involved in alteration of chromatin structure. The rs3803662 TOX3 polymorphism is associated with an increased risk of and overall survival in breast cancer.</td>
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<td>20</td>
<td>VEGF, vascular endothelial growth factor</td>
<td>VEGF plays an important role in tumor angiogenesis. Genetic variation in VEGF may contribute to cancer susceptibility. An association of the CC, CT, or TT genotypes exhibited modification of breast and ovarian cancer risks. The 936_C&gt;T polymorphism in the VEGF gene has a functional influence on disease risks in BRCA1 carriers (65).</td>
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<td>21</td>
<td>PGR, progesterone receptor</td>
<td>The variant progesterone receptor allele named PROGINS polymorphism (rs1042838) is characterized by a 306-bp Alu insertion into intron 7 and two additional sequence variations in exons 4 and 5 of the PGR gene, Val660Leu and His770His. The presence of one or more PROGINS alleles has an increased risk of developing ovarian and endometrial cancers (66).</td>
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<td>22</td>
<td>KRAS, v-Ki-ras2 Kirsten rat sarcoma viral oncogene homolog</td>
<td>A single amino acid substitution of KRAS, a member of the small GTPase superfamily, is responsible for an activating mutation and results in various malignancies, including lung, pancreas and colorectal cancers. Interestingly, the KRAS-variant at rs61764370 is associated with an increased risk of developing ovarian cancer in HBOC families without other known genetic abnormalities (67).</td>
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Germline mutations in BRCA1 and BRCA2 are associated with an increased risk of HBOC syndrome. Non-BRCA mutations only account for a minority of cases. Members of mutation-negative families are also at increased risk of breast and ovarian cancers. Genome-wide association studies identified other genetic factors that increase the risk of breast and ovarian cancers. In addition to BRCA1 and BRCA2 mutations, putative susceptibility genes for hereditary breast and ovarian cancer include BRCA1/2 function modifier genes and DNA repair modifier genes with high penetrance (CDH1, NBS1, NF1, PTEN, TP53 and STK11), moderate penetrance (ATM, ATR, BRIP1, CHEK2, PALB2, CDK1 and RAD50), and low penetrance (FGFR2, LSP1, MAP3K1, TGFB1, TOX3, VEGF, PGR, KRAS and PARP) (3,68). Although some of the factors were validated, various studies suggest the existence of disease susceptibility differences within ethnic populations.

Hereditary condition is hereditary non-polyposis colorectal cancer (HNPCC) syndrome, also known as ‘Lynch syndrome’. Two manifestations of hereditary ovarian cancer are currently recognized: the HBOC syndrome and the HNPCC syndrome. Lynch syndrome has been defined clinically and genetically and is an autosomal-dominant cancer predisposition syndrome that increases the risk for several forms of malignancy, including colorectal (lifetime cancer risk, 70-80%), endometrial (50-60%), stomach cancer (13-19%), renal pelvis (50-60%), brain, as well as transitional cell carcinoma of the ureters and renal pelvis. Mutations in four MMR genes [MLH1 (mutL homolog 1), MSH2 (mutS homolog 2), MSH6 (mutS homolog 6) and PMS2 (postmeiotic segregation increased 2)] are associated with Lynch syndrome and account for another 10% of hereditary ovarian cancer (26). The MMR genes encode proteins involved in the same pathway of DNA mismatch repair. These genetic defects in the DNA MMR system result in DNA mismatch repair errors, including base substitutions and insertion-deletion loops, known as microsatellite instability (MSI). MutS α complex (composed of MSH2 and MSH6) or MutS β complex (MSH2-MSH3) recognizes single-base mismatches and small insertion-deletion loops, binds to mismatched DNA, and recruits the MutL α complex (MLH1- PMS2), which leads to strand discrimination and removal of the errors and coordinates the remaining steps in MMR (27-29).

Both HNPCC and HBOC associated ovarian cancer develop along distinct genetic pathways (30). Although some deviating reports exist, breast cancer incidence has been found to be elevated in Lynch syndrome patients (31). Furthermore, germline mutations associated with Lynch syndrome has been described in 2-3% of patients diagnosed with endometrial cancer. Among Lynch syndrome-related cancers, endometrial cancer is riskier than colorectal cancer in terms of estimated lifetime cumulative risk (32). The overall 5-year survival rate for endometrial (88 vs. 82%) or ovarian cancer (64 vs. 58%) was not significantly different between patients with endometrial or ovarian cancer that are associated with Lynch syndrome and the controls with sporadic cases (33,34). Although immunohistochemical analysis of tumor tissue proved to be a good pre-screening test before proceeding to germline mutation analysis, the discordant results are sometimes observed between immunohistochemistry and replication error phenotyping. Notwithstanding this limitation, immunohistochemistry may be helpful in the evaluation of women with a likely diagnosis of Lynch syndrome. Immunohistochemistry for DNA MMR showed loss of proteins in 9% of women with synchronous endometrial and ovarian cancer (35).

As shown in Fig. 1, BRCA1 forms a multi-subunit protein complex, which includes DNA damage repair proteins, including FA and MMR proteins. Recent studies showed a functional interaction between FANCl and the MMR complex MutL α, which is essential for establishment of DNA inter-strand cross-links (36). FANCD2 is required for binding between MSH2 and MLH1, which are involved in the mono-ubiquitination of FANCD2, leading to recruitment of ATR and then activation of CHK1 and TP53. Furthermore, MutS α and MutL α complex have been shown to be required for the
recruitment of ATR to DNA damage lesion. Taken together, these results support that there might be a functional overlap between the MMR and FA-BRCA pathways.

6. Prevention

Attention has been paid to the role of modifiable risk factors like reproductive histories and exogenous hormones. Potential modifying factors include age of menarche, parity, breastfeeding and oophorectomy. Oral contraceptives (OCs) have a significant protective effect on the risk of ovarian cancer by ~50% in the general population (28). OCs reduce the risk of ovarian cancer also in BRCA1/BRCA2 mutation carriers. The effect of parity may be different in BRCA1 and BRCA2 carriers. Parity protects against breast cancer in BRCA1 mutation carriers. Multiparity may be associated with an increase in risk in BRCA2 carriers. However, the effect of multiparity on ovarian cancer risk for BRCA2 mutation carriers has only been investigated in a small number of studies. Therefore, the association is controversial. Women who took OCs before the age of 30 years and long-term user of 5 or more years have been associated with a slight increase in risk of breast cancer among BRCA1 mutation carriers (29). Furthermore, prolonged hormone replacement therapy (HRT) use is an established risk factor of breast cancer. HRT also affects ovarian cancer risk. Among a subgroup of patients who had a familial history of breast cancer, tamoxifen and raloxifene, selective estrogen receptor modulators, reduced breast cancer risk. Tamoxifen and raloxifene reduced the risk of invasive breast cancer: compared with placebo, raloxifene reduced breast cancer risk by 38%, tamoxifen showed a 50% reduction (37). Additionally, use of PARP inhibitors is a potential synthetic lethal therapeutic strategy and may be considered as targeted chemoprevention in patients with specific DNA-repair defects. Other agents under preclinical and clinical investigation include cyclooxygenase-2 inhibitors, aromatase inhibitors, tyrosine kinase inhibitors, and difluoromethylornithine (a polyamine inhibitor) (38,39). Future efficacy studies are expected.

Prophylactic surgeries are appropriate treatment options for BRCA mutation-associated cancer (4). Prophylactic bilateral mastectomy reduces the risk of breast cancer in BRCA mutation carriers by 90% at any age. Prophylactic bilateral salpingo-oophorectomy (BSO) lowers the ovarian cancer risk by 80%. BSO performed before age 50 years also exhibits a 50% reduction in subsequent breast cancer risk. In conclusion, prophylactic surgeries lead to a reduction in breast and ovarian cancer-specific mortality. Knowledge of these risk factors and prevention strategies will have a great impact on the management of hereditary breast and ovarian cancers.

7. Screening

Since identification of the mutation screening is currently labor intensive and expensive, the screening should be directed to asymptomatic individuals only if they belong to high-risk families. Various safe and effective screening protocols have been recommended for early cancer detection and reduction of cancer risk in clinical practice. Women with HBOC syndrome often utilize the latest medical advances in increased surveillance, prevention, early detection, chemoprevention and optimal treatment. Among BRCA1 and BRCA2 mutation carriers, use of screening mammography alone led to increased early detection rates of non-palpable breast cancer,
but the rate of interval cancers was high (7,40). Therefore, the effectiveness of mammography alone (sensitivity 40%) is questionable for screening high-risk women. In the high risk group of women, magnetic resonance imaging (unenhanced MR imaging with combined diffusion-weighted and T2-weighted images, sensitivity 50%) surveillance largely out performed mammography. Furthermore, dynamic contrast-enhanced MRI exhibited highest sensitivity (86%) (41). MRI is a better screening method and will detect the majority of breast cancers at an early stage. The addition of MRI to screening mammography increased sensitivity (42), supporting the benefits of breast MRI examination annually (the sensitivity was 80%, the false positive rate was 10%) in BRCA mutation carriers (43). Alternating MRI and mammography screening at 6-month intervals might be a clinically effective approach. The HBOC carriers have two options to reduce their risk of ovarian cancer: periodic screening and risk-reducing surgeries (44). Periodic screening consists of annual or semi-annual pelvic examination with the longitudinal CA125 blood test plus concurrent transvaginal ultrasound. Unfortunately, to date, this screening regimen is ineffective for early detection of ovarian cancer in high-risk women. Risk-reducing salpingo-oophorectomy represents a potentially valuable intervention and is the only way for many women at high-risk by age 40 years, or on completion of childbearing. Preventive surgery can reduce ovarian cancer risk by 80-90% and breast cancer risk by 50-60% in BRCA mutation carriers.

8. Discussion

HBOC syndrome is the inherited tendency to develop breast, ovarian and other cancers and believed to be transmitted by mutations in the specific genes. Clinical characteristics, including the type of tumor and age at occurrence as well as family history, predict the prevalence of BRCA germline mutations. A number of clinicians usually take into account the age of the youngest breast cancer patient and the number of ovarian cancer cases in a family as well as pathological diagnosis. Up to 80% of the HBOC cases are due to mutations in BRCA1 or BRCA2 genes. Both BRCA1 and BRCA2 mutations are scattered throughout the whole coding exons.

To maintain and restore the genomic integrity, normal cells possess DNA repair mechanisms. The structural modifications, such as DNA base damage, DNA strand break, inter- and intra-strand crosslinks and DNA-protein crosslinks, are involved in mutation and cancer. A variety of intelligent mechanisms can activate DNA repair pathways and cell cycle checkpoints and recognize and repair SSB or DSB by the master sensors and regulators of DNA damage response such as BRCA1 and BRCA2. BRCA1 and BRCA2 genes were recruited to the sites of DNA damage. BRCA1 associates with several proteins and is an integral member of the repair of DNA damage by functional HR, NER and possibly NHEJ. BRCA1 activated by edk1 physically interacts with MutLα, through the interaction of the FA-BRCA pathway (45). BRCA2, also known as FANC2D, has a more specific role in DNA repair and is directly involved in the mechanism of HR, regulating the activity of RAD51, a gene implicated in the HR pathway as well as interacting with PALB2, a gene implicated in the HR repair and checkpoint functions. Interestingly, the nuclear function of BRCA proteins is tightly regulated by the FA-BRCA, MMR, BER and NER pathway. However, the exact mechanisms of the BRCA-associated interaction and accumulation of other DNA repair proteins are not comprehensively known. Therefore, germline mutations in other susceptibility genes, such as FA genes, MMR genes and DNA repair genes, might be the predisposing factors in HBOC cases. These predisposing genes encode for upstream and downstream regulators of BRCA gene products and also may be associated with the BRCA core complex, including mutations in FANCD1, FANCD2, FANCJ, FANCN, TP53, PTEN, STK11, CDH1, CHK2, ATM, ATR, MSH1, M SH2, MLH1 and PMS2 genes. Families affected by other syndromes, such as Lynch syndrome (mutations in MMR genes), Fanconi anemia, Cowden syndrome (mutations in PTEN), Li-Fraumeni syndrome (mutations in TP53), xeroderma pigmentosum and ataxia-telangiectasia, exhibit additional types of cancers outside the previously defined HBOC cancer spectrum.

In conclusion, genetic or epigenetic loss-of-function mutations of genes that are known to be involved in the repair of DNA damage might lead to increased risk of developing a broad spectrum of breast and ovarian cancers.

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