A Case of Male Breast Cancer Patient with CHEK2*1100delC Mutation

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Abstract

Male breast cancer (MBC) is a rare disease that accounts for less than one percent of all breast cancers. The association between BRCA1 and BRCA2 mutations and MBC has been well-established; recent data suggest that CHEK2 1100delC heterozygosity is also associated with an increased risk of MBC. Herein, we present the case of a 47-year-old male who was initially diagnosed with bilateral symmetric gynecomastia on a diagnostic mammogram performed for right breast palpable lump. Sixteen months after his diagnosis of gynecomastia, he presented with enlarging right breast palpable lumps and underwent a diagnostic mammogram and breast ultrasound. Ultrasound-guided biopsies were performed on the right breast mass and axillary lymphadenopathy. Pathology revealed right breast invasive ductal carcinoma (IDC) and right axillary metastatic lymphadenopathy. Subsequent genetic testing found CHEK2*1100delC mutation. This case report focuses on the presentation, diagnosis, and management of breast cancer, as well as long-term cancer screening in the setting of CHEK2 mutation in a relatively young male patient.

Introduction

Male breast cancer (MBC) is a rare occurrence, comprising less than 1% of cancer in men and less than 0.2% of all cancer-related mortality among men. Risk factors for the development of breast cancer in men include alcohol use, as well as factors that could lead to a hyperestrogenic state, such as obesity, cirrhosis, testicular injury, and undescended testes [1]. Moreover, several breast cancer susceptibility genes are known risk factors for breast cancer, including BRCA1 and BRCA2 mutations. In addition, CHEK2 mutations have been shown to predispose both women and men to specific types of breast cancer. CHEK2 protein is a cell-cycle checkpoint kinase, responsible for phosphorylating the tumor suppressor proteins p53 and BRCA1, which subsequently leads to either cell cycle arrest or activation of DNA repair proteins. Loss-of-function variants in CHEK2 are known to be pathogenic as they result in impaired DNA repair and genomic instability. More specifically, the 1100delC variant results in the deletion of one nucleotide from exon 11 of the CHEK2 mRNA, causing a frameshift at codon 367; this forms a premature stop codon, creating a disrupted or absent protein product and abolishing the kinase function of CHEK2 [2]. CHEK2 mutation is inherited in an autosomal dominant manner [3].

The association between CHEK2 mutation and female breast cancer has been extensively studied in the past. According to one meta-analysis by Weischer et al., CHEK2 1100delC variant can increase the risk of breast cancer three- to five-fold among women; in addition, women with a family history of breast cancer are estimated to have a 37% cumulative risk of breast cancer at age 70 years, while those with no family history of breast cancer are estimated to have a 21% cumulative risk [4]. Another meta-analysis by Yang et al. confirmed the significant association between the CHEK2 1100delC variant and female breast cancer risk, particularly in familial breast cancer cases among Caucasians [5].

CHEK2 mutation has also been associated with an increased risk of MBC, although the association has not been as well-studied as in female breast cancer [6]. From the articles that have studied this association between CHEK2 mutation and MBC thus far, however, one can conclude that the prevalence of CHEK2 mutation is higher in some countries compared to the others. Higher rates of CHEK2 mutation have been reported in studies from Northern European countries, mainly the Netherlands; the mutation, however, seems to be rare in Australia, Spain, and Ashkenazi Jews [7]. Therefore, the evaluation of the association between CHEK2 mutation and MBC has been limited so far due to both the rarity of the MBC cases as well as the significant differences in the CHEK2*1100delC population frequencies [6]. Herein, we present a case of MBC with CHEK2 mutation.

Case Presentation

The patient is a 47-year-old male with a past medical history of bilateral gynecomastia that was first
detected via a mammogram in November 2018 with no evidence of malignancy present on mammogram at the time (Figure 1). In addition, he has a past surgical history of Roux-en-Y gastric bypass in 2007 with revision done in 2013, with his most recent body mass index being 37 kg/m². His social history is positive for a 45-pack-year history of cigarette smoking, which he has now quit, as well as heavy alcohol abuse until one year ago, which he has recently cut down to one shot of whiskey a day. Sixteen months after his diagnosis of gynecomastia, the patient presented with a two-month history of right retroareolar palpable mass along with new-onset tingling, numbness, and pain of the right breast. On exam, the patient had a right-sided subareolar mass measuring 5 x 3 cm that was mobile from chest wall but fixed to skin, along with color changes of the nipple. In addition, he had a right axillary mass that was mobile and approximately 3 cm in size. A diagnostic mammogram and breast ultrasound showed an irregular right retroareolar mass along with an abnormal-appearing right axillary lymph node, both of which were highly suggestive of malignancy (BI-RADS 5) (Figures 2, 3). The breast mass measured 35 x 26 x 24 mm and the axillary lymph node measured 19 x 17 x 18 mm on ultrasound (Figures 4, 5). Ultrasound-guided core biopsy of the right-sided retroareolar was performed with pathology revealing cT2, cN1, cM0, Grade 3, ER/PR positive, HER2 negative, Clinical Stage IIB invasive ductal carcinoma of the right breast. Additional metastatic workup imaging with computed tomography (CT) scan of chest, abdomen, and pelvis was negative (Figure 6).
FIGURE 3: Right breast diagnostic mammogram spot compression CC and LM Views

Right breast irregular mass measuring 35 mm (red circle); craniocaudal (left image) and lateromedial (right image) views.

FIGURE 4: Right breast diagnostic ultrasound

Right breast irregular mass with internal vascular color flow measuring 35 x 26 x 24 mm (red circle).
The patient underwent a modified radical mastectomy with sentinel lymph node mapping; post-mastectomy pathology report confirmed pT2, pN1a, M0, Grade 3, ER/PR positive, Ki-67 50%, HER2 FISH not amplified, Path Stage IIA invasive ductal carcinoma, with negative surgical margins. The patient is currently followed by medical oncology with the plan for Adriamycin/Cyclophosphamide-Taxol (AC-T) regimen with curative
In addition to breast cancer, five years of Tamoxifen treatment.

scheduled to undergo a course of AC-T chemotherapy and receive post-surgical radiotherapy and at least
intraductal carcinoma and positive axillary lymph node, underwent a modified radical mastectomy and is

Adjuvant chemotherapy is also associated with improved survival and is recommended in male patients with

[16] the indications for post-surgical radiotherapy in MBC are the same as for female breast cancer
[15] important part of the surgical treatment of invasive MBC is axillary lymph node dissection
[14] recurrence or survival. Breast-conserving surgery (lumpectomy) is another option but is usually not feasible
[13] due to the rarity of breast cancer in men as well as a lower index of suspicion at initial
[12] Male patients presenting with breast symptoms such as breast enlargement, nipple discharge, breast pain,
[11] The risk factors for breast cancer development in our patient included obesity and heavy alcohol use, in addition to CHEK2 mutation; the combination of all these risk factors could have led to a relatively early onset of breast cancer in this patient.

Although not as extensively studied as in female breast cancer, recent studies have explored the link
between CHEK2 mutation and MBC. According to a study by Wasilewski et al., CHEK2 1100delC is
associated with an increased risk for MBC particularly in the Dutch population, with the average age at
diagnosis of 69 years; most of the CHEK2 1100delC MBC cases in the study were also ER/PR positive
compared to CHEK2 1100delC negative cases [6]. Another study by Cybulski et al. investigated the link
between CHEK2 mutation and ER status in women with early-onset breast cancer and found a four-fold
increased risk of ER-positive breast cancer, suggesting that Tamoxifen could be used as chemoprevention in
the setting of confirmed CHEK2 mutation [12].

According to a study by Marjanka et al., breast tumor characteristics of CHEK2 mutation carriers do not
differ from those of non-carriers, with no significant differences observed in terms of tumor morphology,
angioinvasion, or lymph node involvement [13]. Therefore, the same course of treatment is generally
followed for MBC in CHEK2 carriers as non-carriers. Due to the rarity of MBC, recommendations regarding
its management is based upon the results of clinical trials focusing on female breast cancer. Surgery is
usually the first step in the treatment of MBC. Modified radical mastectomy is generally preferred over
radical or total mastectomy due to a lower rate of complications despite no significant difference in cancer
recurrence or survival. Breast-conserving surgery (lumpectomy) is another option but is usually not feasible
due to the central location of most male breast tumors, as well as the small breast volume in men. Another
important part of the surgical treatment of invasive MBC is axillary lymph node dissection [14]. In addition,
the indications for post-surgical radiotherapy in MBC are the same as for female breast cancer [15]. Adjutant
radiation therapy is associated with improved survival, particularly in patients with positive lymph nodes
[16]. In terms of systemic therapy, men with advanced breast cancer should be managed similarly to women.
Since the majority of male breast cancers are ER-positive, at least 5–10 years of Tamoxifen therapy is
recommended. In MBC, however, if Tamoxifen is contraindicated, a gonadotropin-releasing hormone
(GnRH) analog is used alongside aromatase inhibitors to promote adequate estradiol suppression [15].
Adjutant chemotherapy is also associated with improved survival and is recommended in male patients with
positive axillary lymph node status [14]. The same chemotherapeutic agents that are recommended in
invasive female breast cancer should be used for invasive MBC [11]. Our patient, with ER/PR positive
intraductal carcinoma and positive axillary lymph node, underwent a modified radical mastectomy and is
scheduled to undergo a course of AC-T chemotherapy and receive post-surgical radiotherapy and at least
five years of Tamoxifen treatment.

In addition to breast cancer, CHEK2 mutation can predispose to other types of cancer. According to a study
by Näslund-Koch et al., CHEK2 heterozygosity is associated with a 15–82% increased risk for any cancer; they
propose that this mutation could be a susceptibility allele for any cancer, especially given its function as a protein kinase involved in DNA repair and stability [17]. Another study by Kleibalova et al. suggests that second primary cancers, such as cancers of colon, thyroid, lung, and kidney, are more frequent in CHEK2 mutation carriers compared to non-carriers [18]. In addition, two meta-analyses, one by Liu et al. and another by Xiang et al., have explored the link between the CHEK2 mutation and colorectal cancer and have found an increased risk for colorectal cancer among CHEK2 mutation carriers [5,19]. For this reason, NCCN guidelines suggest colonoscopy screening every five years beginning at age 40 if there are no first-degree relatives diagnosed with colorectal cancer or every five years beginning at age 40 or 10 years prior to the age of a first-degree relative with colorectal cancer [20]. Our patient was also recommended to schedule a colonoscopy in the future following his oncology treatment.

Moreover, since CHEK2 mutation is inherited in an autosomal dominant fashion, our patient was informed that other family members may also carry this pathogenic variant. He was informed that children of CHEK2 carriers also have a 50% risk to carry the pathogenic variant with a 50% chance for his son to carry the familial CHEK2 pathogenic variant. Since it is unclear whether the CHEK2 pathogenic variant was maternally or paternally inherited at this point, testing his maternal and paternal relatives can help determine which family members are also at risk of carrying the pathogenic variant. The patient was encouraged to discuss the results with his family members and to talk to them regarding seeking genetic counseling to clarify their risk.

Conclusions

In most cases, when a male presents with a breast lump, the diagnosis will be gynecomastia. However, MBC is also on the differential, and thus, assessment with a diagnostic mammogram and breast ultrasound can be helpful. MBC is uncommon, and it typically occurs in elderly men with a peak age of 71 years old. In this case, the patient is only 47 years old. Although he is relatively younger than the peak age for MBC, his genetic predisposition with the CHEK2 mutation explains his diagnosis of MBC at a relatively younger age. The patient also has gynecomastia, but gynecomastia is not a risk factor for breast cancer according to the current literature. Breast cancer genetic testing is important after diagnosis of MBC to provide further information to the patient as well as his family members. Due to the association of CHEK2 mutation with other malignancies, further cancer surveillance is recommended for patients with this mutation.

Additional Information

Disclosures

Human subjects: All authors have confirmed that this study did not involve human participants or tissue.
Conflicts of interest: In compliance with the ICMJE uniform disclosure form, all authors declare the following: Payment/services info: All authors have declared that no financial support was received from any organization for the submitted work. Financial relationships: All authors have declared that they have no financial relationships at present or within the previous three years with any organizations that might have an interest in the submitted work. Other relationships: All authors have declared that there are no other relationships or activities that could appear to have influenced the submitted work.

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