PRACTICE POINTER

Care of men with cancer-predisposing BRCA variants

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What you need to know

- Men and women are equally likely to inherit or pass on a cancer-predisposing BRCA variant—family history of cancers needs to encompass both sides of the family
- Men with cancer-predisposing BRCA variants have an increased risk of developing breast cancer and are advised to be breast aware
- Men with cancer-predisposing BRCA2 variants have an increased risk of developing aggressive prostate cancer (men with cancer-predisposing BRCA1 variants may also have an increased risk); it is not yet known whether prostate specific antigen screening reduces mortality in men with cancer-predisposing BRCA variants
- The European Association of Urology recommends that PSA screening is offered to men with cancer-predisposing BRCA2 variants from 40 years of age after discussion of the risks and benefits

Around one in 260 men (~0.4%) inherits a cancer-predisposing BRCA variant that increases their risk of developing prostate, pancreatic, and breast cancer and may affect the health of their family. 1, 2 Most of these men are currently unaware that they have a cancer-predisposing BRCA variant, but as genetic testing becomes more common, more men will need medical advice about what having such a variant means for them and their families.

Men are just as likely as women to have a cancer-predisposing BRCA variant, but many people perceive these variants as only being relevant to women. Paradoxically, this could lead to women at very high risk of breast and ovarian cancer missing out on screening and risk-lowering treatment despite a concerning paternal family history. Clinicians might also be less attuned to paternal family history of cancer in assessing women’s breast cancer risk. 3 This practice pointer covers what cancer-predisposing BRCA variants are, who might be tested; and what health issues men and their clinicians need to know about. We refer to men, but the article also applies to transwomen and some non-binary people.

What are cancer-predisposing BRCA variants?

BRCA1 and BRCA2 are tumour suppressor genes that code for DNA repair proteins. Certain variants in these genes predispose to cancer (primarily breast, ovarian, prostate, and pancreatic, and for BRCA2 possibly melanoma 4) (fig 1). The predisposition to cancer is inherited in an autosomal dominant way, ie, each and every time a person with a cancer-predisposing BRCA variant has a child, they have a 1 in 2 or 50:50 chance of passing their BRCA variant on to the child. This is regardless of whether parent or child is male or female.

Many men and some women with cancer-predisposing BRCA variants will never develop an associated cancer (fig 1). The cancer risks associated with cancer-predisposing BRCA variants are modified by lifestyle factors and other inherited genetic variants. Polygenic risk scores (calculated by looking at multiple common genetic variants across the genome, each with a tiny individual effect) show potential to refine cancer risk predictions for people with cancer-predisposing BRCA variants, but cannot remove uncertainty as to whether a given patient will or will not develop cancer. 10

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Fig 1(a) | Risk of ovary, prostate, and pancreatic cancer associated with cancer-predisposing BRCA variants (cumulative cancer risks are shown on different scales).
Cancer-predisposing BRCA1 variants increase the risk of pancreatic cancer (relative risk 4.11) but data are emerging, and cumulative risk figures are not readily available. Data on ovary cancer from Kuchenbaecker et al 2017; prostate from Nyberg et al 2020, and pancreatic cancer from van Asperen et al 2005. Population data are for England and Wales 2016 (Office for National Statistics).

Fig 1(b) | Risk of female and male breast cancer associated with cancer-predisposing BRCA variants. Data from Tai et al 2007. Population data are for England and Wales 2016 (Office for National Statistics).

Describing genetic variants
Various terms are used to describe genetic variants that increase risk of disease. As with many medical terms, different people might understand different things by the same term. The term "mutation" is often used in patient leaflets, patient support forums, and in UK school curriculums. However, some find the term offensive and it is technically incorrect since it suggests the mutation has happened in that person. "Faulty gene" is also often used in patient resources, and a 2007 Australian survey indicated that this term was most preferred by the general community and by patients recruited via cancer genetics clinics.
"Pathogenic" or "likely pathogenic variant" is commonly used in laboratory genetic reports. People are increasingly familiar with the term "variant" in the context of covid-19, but "pathogenic" may be less well
understood by people without a healthcare background, so “disease-causing” is sometimes used instead. However, this in turn may give the false impression that everyone with a "disease-causing BRCA variant" will inevitably develop an associated cancer. Other terms in common use include: "disease-risk variant," "genetic risk factor," "disease-predisposing variant," "loss-of-function variant," "gene alteration," "gene change" and for people with cancer-predisposing BRCA variants, "BRCA carriers." Importantly there are also many variants in BRCA1 and BRCA2 which are not associated with cancer risks.

In discussion with our hospital Patient and Public Involvement group it was evident that different people found different terms most helpful. We use the term "cancer-predisposing variant" here as it is informative and technically accurate, but whatever term you use to describe cancer-predisposing BRCA variants, key messages to convey are:

- variants are very likely to be inherited rather than to have happened for the first time in the tested person
- variants are risk factors for particular cancers rather than certainties that they will develop.

Thousands of different variants within the BRCA1 and BRCA2 genes have been described, and classifying which are benign and which increase cancer risk can be challenging. As more data are gathered, classifications may shift. For example a Canadian laboratory examined the BRCA variants they identified over a five year period and found that 12% were reclassified (75% of these were downgraded—ie, they are now thought to be less likely to predispose to cancer than they were when the BRCA test was originally done). Diagnostic tests involve sequencing of BRCA1 and BRCA2 (and perhaps other genes, eg, PALB2), aiming to detect any variant present that might increase cancer risk.

How would a man find out that he has a cancer-predisposing BRCA variant?

In the UK, diagnostic BRCA testing is currently offered to people with a 10% or greater chance of having a cancer-predisposing BRCA variant. Most diagnostic BRCA tests are done for women, but some men will find out their BRCA status via this route, as men with breast cancer at any age are eligible to be tested, and some men with prostate or pancreatic cancer may have testing to determine eligibility for poly(ADP-ribose) polymerase (PARP) inhibitors (often in the context of clinical trials).

More commonly, men are offered targeted BRCA testing after a cancer-predisposing BRCA variant is identified in their family. Usually this will be predictive testing (ie, the man has no personal medical history suggestive of a cancer-predisposing BRCA variant), although occasionally it may be explanatory (eg, in men known to have prostate cancer). The laboratory would need details of the specific cancer-predisposing BRCA variant in the man’s family in order to test for it (eg, a genetic report from an affected family member) and such targeted testing would not detect any other cancer-predisposing variants.

Around 1.5% of patients with prostate cancer have a cancer-predisposing BRCA1 variant, but a personal medical history of prostate cancer is not an indication for diagnostic BRCA testing in the UK. However, for patients with younger-onset, aggressive disease, a thorough family history can explore the possibility of a familial cancer-predisposing BRCA variant. A family history that includes, for example, bilateral breast cancer, male breast cancer, or multiple people affected by breast or ovarian cancer (especially at younger ages), probably warrants discussion with your local genetics service.

Some men might access BRCA testing outside standard clinical pathways, such as through direct-to-consumer genetic testing or research studies. These tests are of variable quality and scope, and further scrutiny may be needed to confirm that the variant is really present or really represents a risk.

Case studies

Cancer-predisposing BRCA variants increase the risk of aggressive prostate cancer

Rahul is a 40 year old man who had a test via a genetics clinic for the cancer-predisposing BRCA2 variant identified in his aunt. Rahul was also found to have the BRCA2 variant and is concerned about developing prostate cancer. He books an appointment with you to discuss prostate specific antigen (PSA) screening.

Men with cancer-predisposing BRCA2 variants like Rahul have an increased risk of prostate cancer: a recent meta-analysis indicated an odds ratio of 2.64, and a large prospective cohort study in the UK and Ireland found a 27% absolute risk of developing prostate cancer by age 75, rising to 60% by age 85 (for comparison, population risk by age 85 is 16% in England and Wales based on Office for National Statistics 2016 data). When prostate cancer occurs in a man with a cancer-predisposing BRCA2 variant, it tends to be more aggressive. The evidence has been inconsistent regarding the impact of cancer-predisposing BRCA1 variants on prostate cancer but a subtler effect is probable, with an odds ratio of 1.35, and the relative risk increase is higher at younger ages.

The IMPACT study is an ongoing international prospective cohort study of more than 3000 men to examine the use of PSA screening in men with cancer-predisposing BRCA variants. Based on interim results, the researchers recommend that men with cancer-predisposing BRCA2 variants are offered systematic PSA screening, and this could be considered for Rahul. This is because after three years of PSA screening, men with cancer-predisposing BRCA2 variants proved to have a higher incidence of prostate cancer, were younger at diagnosis, and were more likely to have clinically significant tumours. The positive predictive value of a PSA >3.0 ng/mL was higher in men with cancer-predisposing BRCA2 variants than in controls (31% versus 18%).

Discuss with Rahul the pros and cons of PSA screening and be clear about the current limits of medical knowledge. It would be appropriate to include in this discussion that if a prostate cancer does develop in a man with a cancer-predisposing BRCA2 variant, it is more likely to be clinically significant. Interim analysis from the IMPACT study shows that after four screening rounds (annual PSA) for men aged 55-69 with cancer-predisposing BRCA2 variants, you would expect to detect one clinically significant prostate cancer for every 13 men screened. The European Association of Urology recommends offering PSA based prostate cancer screening to men with cancer-predisposing BRCA2 variants who have been counselled on the potential risks and benefits of screening from the age of 40 although they do not specify a screening interval. It is not yet known whether PSA screening will reduce mortality from prostate cancer in men with cancer-predisposing BRCA2 variants. Research is ongoing as to the role of PSA screening for men with cancer-predisposing BRCA1 variants: an interim analysis found no differences in age or tumour characteristics between men with cancer-predisposing BRCA1 variants and controls.
Breast awareness is important for men with cancer-predisposing BRCA variants

Jakob is a 50 year old man who books an appointment to discuss a painless “cyst” near his left nipple that he noticed several months ago. On examination, you notice that he has an inverted nipple and ipsilateral axillary lymphadenopathy. When you ask about his family history, he tells you that his father died of prostate cancer in his 60s.

Men with cancer-predisposing BRCA variants have an increased lifetime risk of developing breast cancer: 8.3% for BRCA2 and 1.8% for BRCA1, compared with 0.1% in the general population. As in women, breast cancer in men most commonly presents as a painless mass. Although nipple involvement tends to be seen earlier due to the smaller amount of breast tissue, male breast cancer is often diagnosed at an advanced stage. Men with cancer-predisposing BRCA variants are at higher risk of developing breast cancer, but any man with symptoms of breast cancer warrants urgent referral to a breast clinic in the same way women presenting with concerning symptoms would be referred, regardless of BRCA status.

Evidence regarding breast cancer characteristics in men with cancer-predisposing BRCA variants is limited. Two studies analysing tumour grading, staging, and receptor status in men with breast cancer suggest that cancer-predisposing BRCA2 variants are associated with more aggressive cancers.

Evidence is also lacking on breast cancer screening in men with cancer-predisposing BRCA variants, and practice varies. In the UK, men with cancer-predisposing BRCA variants are advised to be breast aware, ie, to know how their breasts usually look and feel, and seek medical advice if they notice changes or have any concerns. Men might be more likely to delay seeking care for a new breast lump, perhaps waiting until the lump becomes painful or changes the overlying skin. A study in Hong Kong of men with breast cancer found that median duration from symptoms to first medical consultation was 12.4 months, and 84% were not aware (before their diagnosis) that breast cancer could occur in men.

Raising this issue may be challenging, particularly as resources promoting breast awareness are mainly aimed at women. Our hospital Patient and Public Involvement group highlighted some of the issues men might face during and after receiving a breast cancer diagnosis, for example being the only man in the waiting room for appointments.

Patients may be unaware that cancer-predisposing BRCA variants can be passed on by men

Harry is a 55 year old man whom you see regularly for diabetes management. At the end of an appointment discussing his blood sugars, he mentions that his sister in Australia recently told him that she “has BRCA” and that he should get tested. You ask how he feels about this, and he says he can’t see the point because “isn’t BRCA a female thing?” Harry has two daughters in their 30s.

Because cancer-predisposing BRCA1 and BRCA2 variants are notorious for increasing lifetime risk of breast and ovarian cancer in women, there is a common misperception that cancer-predisposing BRCA variants themselves only occur in women. Cancer-predisposing BRCA variants are just as common in men as they are in women but are less likely to be detected because they are less likely to cause a cancer that prompts genetic testing.

Explain that men and women inherit these genetic variants in the same way: that any child of a parent with a cancer-predisposing BRCA variant has a 50:50 chance of inheriting that variant. This would include Harry’s children if Harry also has the BRCA variant identified in his sister.

For some men, concern for the health of existing or potential daughters or granddaughters is a key motivation for seeking BRCA testing. In contrast, sometimes men are reluctant to have BRCA testing because they are concerned about potentially having passed a cancer-predisposing BRCA variant on to their children. They might prefer not to know than to have this possibility confirmed. It may help to remind them that the cancer-predisposing BRCA variant has likely been in their family for generations and whether they inherited or passed it on is outside their control. However, being tested for it might guide medical care for them and their children, for example by informing choices about cancer screening (fig 2).
Occasionally, men decide that they do not wish to be tested for a cancer-predisposing BRCA variant found in their family. It would still be appropriate and important to refer their adult, first degree relatives (eg, siblings or children) to clinical genetics specialists if they wish, even if testing them might (indirectly) reveal that the man has the cancer-predisposing BRCA variant. Clinical genetic services are skilled at counselling patients through difficult psychosocial and ethical issues such as this scenario (box 2).

**Box 2: Ethical issues**

- **Sharing genetic information within families**
  - Family members may benefit from testing for the cancer-predisposing BRCA variant. What if the person in whom it was identified finds it difficult to tell their family about the variant, or chooses not to? Health professionals may sometimes need to balance the competing tensions of patient confidentiality with the interests others have in knowing about their risks.

- **Decisions around termination of pregnancy owing to adult onset conditions**
  - People with cancer-predisposing BRCA variants have a substantially increased risk of developing certain cancers, but these are adult-onset and some people with such variants will never develop cancer. Classifying BRCA variants as benign or cancer-predisposing is technically challenging. Over time, new evidence may shift our understanding of what a particular variant means (eg, it may become clear that a variant thought to predispose to cancer is actually benign, or vice versa). How should clinicians respond when this happens?

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**Men with cancer-predisposing BRCA variants have an increased risk of developing pancreatic cancer**

Simon is a 50 year old man who had a test two years ago via a genetics clinic for the cancer-predisposing BRCA2 variant identified in his sister. Simon was also found to have the BRCA2 variant and discussed his increased risk of prostate and pancreatic cancer with a genetic counsellor at the time. Last week Simon’s friend was diagnosed with pancreatic cancer. Simon is now more concerned about his risk of pancreatic cancer and books an appointment to discuss it with you.

People with cancer-predisposing BRCA variants have an increased risk of developing pancreatic cancer: a retrospective cohort analysis of a high-risk breast cancer family registry from the US, Canada, and Australia estimated that pancreatic cancer risk is five to six times higher than population risk for people with cancer-predisposing BRCA2 variants and around four times higher for people with cancer-predisposing BRCA1 variants. Unfortunately, pancreatic cancer is challenging to screen for and is often advanced by the time symptoms develop. Guidance from the National Institute for Health and Care Excellence (2018) on pancreatic cancer recommended that people with cancer-predisposing BRCA variants and one or more first degree relatives with pancreatic cancer have surveillance (magnetic resonance imaging/magnetic resonance cholangiopancreatography or endoscopic ultrasound), but this was challenged by the UK Cancer Genetics Group as being premature. Instead, the group recommended that pancreatic cancer screening should only be offered within the context of research studies such as EUROPAC. In practice, pancreatic cancer screening is generally considered on a case-by-case basis for people with cancer-predisposing BRCA variants if they also have a family...
history of pancreatic cancer. Screening as part of a research study might be a possibility for Simon—your local genetics service may be able to signpost towards this.

Around 6-7% of people with metastatic pancreatic cancer have a germline cancer-predisposing BRCA variant. Increasingly, people with pancreatic cancer are offered diagnostic BRCA testing to inform treatment plans. Cancer cells with cancer-predisposing BRCA variants already have impaired DNA repair and are heavily reliant on DNA repair pathways involving PARP, so are particularly vulnerable to PARP inhibitors. PARP inhibitors may be a treatment strategy for men with a germline cancer-predisposing BRCA variant who have breast, ovarian, pancreatic, or prostate cancer (often in the context of clinical trials).

**Education into practice**

- Do you ask about paternal as well as maternal family history when assessing a patient’s risk of breast cancer?
- What questions might you ask if a man tells you he doesn’t want testing for a cancer-predisposing BRCA variant found in his family?
- When did you last ask a man about his family history of breast and ovarian cancer?

**How patients were involved in the creation of this article**

We spoke with men from the Patient and Public Involvement group at University Hospitals Southampton NHS Foundation Trust to discuss what prior knowledge men might have about cancer-predisposing BRCA variants, what information they might want to know after finding that they had a cancer-predisposing BRCA variant, and what terminology they might prefer when talking about genetic variants. These discussions particularly influenced the case study section, the discussion about expectations in “How would a man find out that he has a cancer-predisposing BRCA variant?” and the box “Describing genetic variants.”

**How this article was made**

We developed fictitious cases to illustrate common issues that may arise for men with cancer-predisposing BRCA variants, based on scenarios encountered by our regional genetics department.

We used PubMed and author research paper archives to search for information about cancer risks and medical care for men with cancer-predisposing BRCA variants. We aimed to quote cancer risks established by large prospective cohort studies. Medical care of men with cancer-predisposing BRCA variants is under-researched and often guidelines are lacking—in these cases, we have drawn on our experience within a UK regional genetics department to aim to reflect UK standard practice.

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Contributorship and guarantor: RH drafted the article, PP edited the draft, advised on selection of evidence and created figure 1. RH edited the draft from her perspective as a GP. AL developed the article, facilitated patient and public input and edited the draft as a whole. AL is the guarantor.

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