

NATIONAL INSTITUTE FOR HEALTH AND CARE EXCELLENCE

Scope for guideline update (starting 2025)

Familial breast cancer: classification, care and managing breast cancer and related risks in people with a family history of breast cancer

NICE is updating its guideline on [Familial breast cancer: classification, care and managing breast cancer and related risks in people with a family history of breast cancer](#) (CG164).

The update will be developed using the methods and processes in [developing NICE guidelines: the manual](#).

Who the guideline update covers

The update will expand the guideline population to cover people who do not have a family history of breast cancer but are suspected of having an inherited predisposition to breast cancer for other reasons. At publication of the update the guideline will be renamed to reflect this wider population.

- For all aspects of the guideline:
 - people aged 18 years and older with a personal history of breast cancer who:
 - ◇ may have an inherited predisposition to breast cancer because of a family history of breast or ovarian cancer or a related cancer, ancestry with a high prevalence of BRCA1 or BRCA2 mutations or because they have features associated with heritable breast cancer
 - ◇ have a confirmed mutation that predisposes them to developing breast cancer.
 - people aged 18 years and over with no personal history of breast cancer who may have an inherited predisposition to be at increased risk of

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1 developing breast cancer because of a family history of breast, ovarian
2 or a related cancer, ancestry with a high prevalence of BRCA1 or
3 BRCA2 mutations or have a confirmed mutation that predisposes them
4 to developing breast cancer who are:

- 5 ◇ women, trans men and non-binary people born with female
6 reproductive organs
- 7 ◇ trans women and non-binary people born with male reproductive
8 organs who have been on gender affirming hormone therapy for 5
9 years or more.

10 • For the consideration of referral for genetic testing only:

- 11 – people aged 18 and over with no personal history of breast cancer who
12 may be at increased risk of developing breast cancer because of a
13 family history of breast, ovarian or related cancer or ancestry with a high
14 prevalence of BRCA1 or BRCA2 mutations, who are:

- 15 ◇ men
- 16 ◇ trans women and non-binary people born with male reproductive
17 organs who have not been on gender affirming hormone therapy for 5
18 years or more.

19 Equality considerations

20 A new [equality and health inequalities assessment impact](#) has been
21 completed.

22 Settings

23 This guideline will cover all health, mental health and social care settings in
24 which NHS care or social care is received or commissioned.

25 Activities, services or aspects of care covered by the 26 guideline update

27 We will consider making recommendations or updating existing
28 recommendations in the following sections of the current guideline:

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1 **Clinical significance of a family history of breast cancer**

- 2 • Family history and carrier probability

3 **Genetic testing**

- 4 • Carrier probability at which genetic testing should be offered
- 5 • Mutation tests
- 6 • Genetic testing for BRCA1, BRCA2 and TP53 mutations within 4 weeks of
- 7 diagnosis of breast cancer.

8 **Surveillance and strategies for early detection of breast** 9 **cancer**

- 10 • Surveillance for women with no personal history of breast cancer

11 **Risk reduction and treatment strategies**

- 12 • Risk-reducing breast surgery for women with or without a personal history
- 13 of breast cancer
- 14 • Hormone replacement therapy.

15 The headings of the sections may change as part of the update and new
16 ones may be introduced.

17 **This guideline update will not cover**

18 We will not be reviewing the evidence on the following:

19 **Clinical significance of a family history of breast cancer**

- 20 • Family history-taking and initial assessment in primary care
- 21 • Family history-taking in secondary care
- 22 • Family history-taking in a specialist genetic clinic
- 23 • Communicating cancer risk and carrier probability.

1 **Information and support**

2 **Care of people in primary care**

- 3 • Care and management of people in primary care
- 4 • Referral from primary care
- 5 • Patient education and information
- 6 • Support for primary care.

7 **Care of people in secondary care and specialist genetic** 8 **clinics**

- 9 • Care and management approach in secondary care
- 10 • Referral to a specialist genetic clinic
- 11 • Care of people in a specialist genetic clinic
- 12 • Genetic counselling for people with no personal history of breast cancer.

13 **Surveillance and strategies for early detection of breast** 14 **cancer**

- 15 • Surveillance for women with a personal and family history of breast cancer
- 16 • Recommendations for all women having surveillance.

17 **Risk reduction and treatment strategies**

- 18 • Risk factors
- 19 • Menstrual and reproductive factors
- 20 • Hormonal contraceptives
- 21 • Breastfeeding
- 22 • Alcohol consumption
- 23 • Smoking
- 24 • Weight and physical activity
- 25 • Chemoprevention for women with no personal history of breast cancer
- 26 • Risk-reducing oophorectomy.

1 We plan to retain the recommendations in these areas, although they may be
2 revised to update language, reflect current policy or practice, and to ensure
3 consistency with new content.

4 **Draft review questions**

5 We have identified the following draft review questions. These may change
6 during guideline development, but the areas covered will remain as listed in
7 the final scope.

8 The areas covered and draft questions will be used to develop more detailed
9 review questions.

10 **Assessing carrier probability**

11 1. What are the optimal methods of assessing the probability of having a
12 pathogenic variant in breast cancer predisposition genes in people who
13 have either:

- 14 • a personal history of breast cancer that is suspected to be
15 heritable or
- 16 • no personal history of breast cancer but a family history of
17 breast, ovarian or a related cancer, or ancestry with a high
18 prevalence of BRCA1 or BRCA2 mutations?

19 **Assessing the risk of developing breast cancer**

20 2. Which risk prediction tools can most accurately predict the risk of
21 developing future breast cancer in either:

- 22 • people with a personal history of breast cancer that is suspected
23 to be heritable or
- 24 • people with no personal history of breast cancer with a family
25 history of breast, ovarian or a related cancer, or ancestry with a
26 high prevalence of BRCA1 or BRCA2 mutations?

Determining the carrier probability at which genetic testing should be offered

3. At what carrier probability should people with breast cancer who have features associated with heritable breast cancer as well as or in addition to a family history of breast, ovarian or a related cancer or ancestry with a high prevalence of BRCA1 or BRCA2 mutations be offered genetic testing?
4. At what carrier probability should people without a personal history of breast cancer, but with a family history of cancer suggestive of heritable breast cancer or ancestry with a high prevalence of BRCA1 or BRCA2 mutations be offered genetic testing?

Surveillance and strategies for early detection of breast cancer

5. At what frequency should people aged 50 to 59 years at moderate risk of breast cancer receive surveillance with mammography?

Risk-reducing surgery

6. How effective and cost effective is risk-reducing mastectomy in people born with female reproductive organs who are at increased risk of heritable breast cancer (considering risk thresholds, previous cancer history, specific gene mutations, and age)?

Economic aspects

We will take economic aspects into account when making recommendations. For each review question (or key areas in the scope), we will review the economic evidence and, where appropriate, carry out economic modelling and analyses, using an NHS and personal social services perspective.

NICE guidance and quality standards that may be affected by this guideline update

- [Ovarian cancer: identifying and managing familial and genetic risk](#) (2024). NICE guideline NG241
- [Early and locally advanced breast cancer: diagnosis and management](#) (2018). NICE guideline NG101
- [Menopause: identification and management](#) (2015). NICE guideline NG23
- [Ovarian cancer](#) (2025). NICE quality standard 18
- [Breast Cancer](#) (2011). NICE quality standard 12

Further information

The guideline update is expected to be published in April 2027.

To follow the progress of the update, see the [guideline in development page](#).

Our website has information about [how NICE guidelines are developed](#).

NICE guidelines cover health and care in England. Decisions on how they apply in other UK countries are made by ministers in the [Welsh Government](#), [Scottish Government](#) and [Northern Ireland Executive](#).

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