The National Institute for Health and Care Excellence (NICE) is producing guidance on using tumour profiling tests (EndoPredict, MammaPrint, Oncotype DX Breast Recurrence Score, Prosigna and IHC4+C) to guide adjuvant chemotherapy decisions in people with early breast cancer in the NHS in England. The diagnostics advisory committee has considered the evidence base and the views of clinical and patient experts.

This document has been prepared for public consultation. It summarises the evidence and views that have been considered, and sets out the draft recommendations made by the committee. NICE invites comments from registered stakeholders, healthcare professionals and the public. This document should be read along with the evidence base (the diagnostics assessment report and the diagnostics assessment report addendum).

The advisory committee is interested in receiving comments on the following:

- Has all of the relevant evidence been taken into account?
- Are the summaries of clinical and cost effectiveness reasonable interpretations of the evidence?
- Are the provisional recommendations sound, and a suitable basis for guidance to the NHS?

Equality issues

NICE is committed to promoting equality of opportunity, eliminating unlawful discrimination and fostering good relations between people with particular protected characteristics and others. Please let us know if you think that the preliminary recommendations may need changing in order to meet these aims. In particular, please tell us if the preliminary recommendations:
Draft recommendations

1.1 EndoPredict, Oncotype DX Breast Recurrence Score and Prosigna are recommended as options for guiding adjuvant chemotherapy decisions for people with oestrogen receptor (ER)-positive, human epidermal growth factor receptor 2 (HER2)-negative and lymph node (LN)-negative early breast cancer, only if:

- they have an intermediate risk of distant recurrence using a validated tool such as PREDICT or the Nottingham Prognostic Index
- information provided by the test would help them and their clinicians make a shared decision on whether or not to have adjuvant chemotherapy
• the companies provide the tests to the NHS with the discounts agreed in the access proposals and
• clinicians and companies make timely, complete and linkable record-level test data available to the National Cancer Registration and Analysis Service under a data collection agreement with NICE (see section 5.24).

1.2 MammaPrint is not recommended for guiding adjuvant chemotherapy decisions for people with ER-positive, HER2-negative and LN-negative early breast cancer because it is not cost effective.

1.3 IHC4+C is not recommended for guiding adjuvant chemotherapy decisions for people with ER-positive, HER2-negative and LN-negative early breast cancer because the analytical validity of the test is uncertain.

2 Clinical need and practice

The problem addressed

2.1 The tumour profiling tests EndoPredict, MammaPrint, Oncotype DX Breast Recurrence Score, Prosigna and IHC4+C provide information on the activity of genes in tumour samples from people with early breast cancer. The results provide a risk profile of a person’s breast cancer, which can be used with other routinely assessed clinical risk factors, such as nodal status and tumour size. It is claimed that the risk profile can be used to better predict the risk of disease recurrence. Some tests also claim to predict relative treatment effects for chemotherapy. This information is intended to help decision-making about adjuvant chemotherapy use.

2.2 It is also claimed that the tumour profiling tests may improve the identification of early breast cancer in people who may not benefit
from adjuvant chemotherapy because there is a low risk of disease recurrence. For these people unnecessary treatment could be avoided, and therefore the comorbidities and negative effects of chemotherapy on quality of life. Also, for people with early breast cancer at low risk of disease recurrence based on clinical and pathological features, the tests could confirm whether their risk is correct. If reclassified as being at high risk of recurrence, those people may benefit from chemotherapy. People with breast cancer and clinicians may also have more confidence that the treatment they are having or recommending is appropriate.

2.3 This assessment evaluates the clinical and cost effectiveness of EndoPredict, MammaPrint, Oncotype DX Breast Recurrence Score, Prosigna and IHC4+C when used to guide adjuvant chemotherapy decisions. The population was people with oestrogen receptor (ER)-positive (or progesterone receptor-positive or both), human epidermal growth factor receptor 2 (HER2)-negative early breast cancer (stages 1 or 2) with 0 to 3 positive lymph nodes.

2.4 This is a full update of NICE’s diagnostics guidance 10 on gene expression profiling and expanded immunohistochemistry tests for guiding adjuvant chemotherapy decisions in early breast cancer management: MammaPrint, Oncotype DX, IHC4 and Mammostrat, which was published in 2013. This recommended Oncotype DX as an option for guiding adjuvant chemotherapy decisions for people with ER-positive, HER2-negative and lymph node-negative early breast cancer if the person was assessed as being at intermediate risk and the company provided Oncotype DX to NHS organisations according to the confidential arrangement agreed with NICE. The guidance also encouraged data collection on the use of Oncotype DX in the NHS, and further research on MammaPrint, IHC4 and Mammostrat. Since publication of the original guidance,
Mammostrat is no longer available and a new test, EndoPredict, has become available.

The condition

2.5 Breast cancer is the most common cancer and the third most common cause of UK cancer-related deaths. One in 8 women and 1 in 870 men will be diagnosed with breast cancer during their lifetime (Cancer Research UK 2016). In 2014, 46,085 women and 332 men were newly diagnosed with breast cancer in England (Office for National Statistics 2016). Most breast cancer develops in women who are over the age of 50 (Cancer Research UK 2016).

2.6 Breast cancer survival depends on the stage of the disease at diagnosis, the treatment received and the biology of the tumour. More than 90% of women diagnosed with early breast cancer survive for at least 5 years, and 78% survive for 10 years (Cancer Research UK 2016). In contrast, only 13% of those diagnosed with advanced disease survive for more than 5 years.

The diagnostics and care pathways

Diagnosis

2.7 Breast cancer may be diagnosed following an abnormal result in the NHS breast cancer screening programme, or after referral for further investigation because of signs or symptoms that could be associated with breast cancer. The referral criteria are described in NICE’s guideline on suspected cancer.

2.8 When cancer cells have been detected in a biopsy sample, further tests are done to provide more information on the characteristics of the tumour. The results of these tests are used to categorise breast cancer into molecular subtypes and determine which types of treatment it is most likely to respond to. Recommendations on tumour testing are in NICE’s guideline on early and locally...
advanced breast cancer. This guideline is being updated. Tumour tests can include hormone receptor and HER2 tests. Although not routinely done, some laboratories may also test for Ki67, a marker of cell proliferation.

Care

2.9 NICE’s guideline on early and locally advanced breast cancer describes the care pathway. Surgery is often the initial treatment. Neoadjuvant treatment may be used before surgery, to reduce the size of the tumour and enable breast-conserving surgery.

2.10 After surgery, further treatment (adjuvant treatment) may be needed and this can include radiotherapy, chemotherapy, hormone therapy, biological therapy or a combination of these. The decision to offer adjuvant therapy, and the treatments to use, is made taking into account the clinical history, the stage of disease, the likely course of the disease (prognosis), the molecular characteristics of the tumour and the person’s preferences.

2.11 A variety of tools are available that can help to predict the likelihood of breast cancer recurrence based on clinical and pathological features. These may be used to provide prognostic information for patients and to guide the selection of adjuvant therapy. Expert advice suggests that the PREDICT tool version 2.0, an online prognostic and treatment benefit calculator, is the most widely used tool in the NHS in England to calculate risk of recurrence. Adjuvant! Online is not currently available to the NHS.

3 The diagnostic tests

3.1 The assessment compared 5 intervention tests with 1 comparator.
The interventions

EndoPredict (Myriad Genetics)

3.2 EndoPredict is a CE-marked assay that is designed to predict the likelihood of metastases developing within 10 years of an initial breast cancer diagnosis. The test is for pre- and postmenopausal people with early breast cancer with oestrogen receptor (ER)-positive, human epidermal growth factor 2 (HER2)-negative and lymph node (LN)-negative or LN-positive disease (up to 3 positive nodes).

3.3 EndoPredict measures the expression of 12 genes: 3 proliferation-associated genes, 5 hormone receptor-associated genes, 3 reference (normalisation) genes and 1 control gene.

3.4 EndoPredict needs RNA extracted from a formalin-fixed, paraffin-embedded (FFPE) breast cancer tissue sample. The test can be done in a local laboratory or the Myriad Genetics pathology laboratory in Germany. It takes approximately 2 days to get the results from a local laboratory, and longer if samples are sent to Germany.

3.5 The test involves a reverse transcription-quantitative polymerase chain reaction. Online evaluation software calculates an EP score and an EPclin score. An EP score of 0 to less than 5 indicates low risk of distant disease recurrence in the next 10 years. An EP score of 5 to 15 indicates high risk of distant disease recurrence in the next 10 years.

3.6 The EPclin score estimates the probability of metastases developing within 10 years (assuming 5 years of endocrine therapy). It is calculated by adding clinical data about tumour size and nodal status to the EP score. An EPclin score of less than 3.3 indicates low risk (less than 10%) of metastases in the next
10 years. An EPclin score of 3.3 or more indicates high risk of metastases in the next 10 years.

3.7 During consultation on the first diagnostics consultation document, NICE accepted an access proposal from the company in line with the Diagnostics Assessment Programme’s interim addendum on access proposals. This provides a simple discount to the list price of EndoPredict, with the discount applied at the point of purchase or invoice. The level of the discount is commercial in confidence.

**MammaPrint (Agendia)**

3.8 MammaPrint is a CE-marked assay that is designed to assess the risk of distant recurrence within 5 and 10 years and whether a person would benefit from chemotherapy. The test is for pre- and postmenopausal people with stage 1 or 2 breast cancer, with a tumour size of 5 cm or less, and LN-negative or LN-positive disease (up to 3 positive nodes). The test can be used irrespective of ER and HER2 status.

3.9 MammaPrint measures the expression of 70 genes, including genes associated with 7 different parts of the metastatic pathway: growth and proliferation, angiogenesis, local invasion, entering the circulation, survival in the circulation, entering organs from the circulation, and adaption to the microenvironment at a secondary site.

3.10 The MammaPrint test needs RNA extracted from an FFPE breast cancer tissue sample. The test is offered as an off-site service. In Europe, samples are analysed at the Agendia laboratory in the Netherlands. Results are available within 10 days of submitting the sample.

3.11 The test is based on diagnostic microarray. Software is used to calculate the MammaPrint result on a scale of −1 to +1. The score
indicates the risk of developing distant metastases over the next 10 years without any adjuvant endocrine therapy or chemotherapy. A MammaPrint result of 0 or less indicates high risk of metastases in the next 10 years and a result of more than 0 indicates low risk (10% or less) of metastases in the next 10 years.

**Oncotype DX Breast Recurrence Score (Genomic Health)**

3.12 Oncotype DX Breast Recurrence Score (hereafter referred to as Oncotype DX) is designed to quantify the 10-year risk of distant recurrence and predict relative treatment effects for chemotherapy. The test also reports the underlying tumour biology: ER, progesterone receptor and HER2 status. The test is for pre- and postmenopausal people with stage 1 or 2 breast cancer and ER-positive, HER2-negative, LN-negative or LN-positive disease (up to 3 positive nodes). The assay does not have a CE mark because it is provided as a service by Genomic Health.

3.13 Oncotype DX quantifies the expression of 21 genes: 16 cancer-related genes correlated with distant recurrence-free survival, and 5 reference (normalisation) genes.

3.14 The Oncotype DX test needs RNA extracted from a FFPE breast cancer tissue sample. Samples are processed centrally at a Genomic Health laboratory in the US. Results are usually available 7 to 10 days after the sample is received.

3.15 The test is based on a reverse transcription-quantitative polymerase chain reaction. It gives a recurrence score of between 0 and 100, which is used to quantify the 10-year risk of distant recurrence, assuming 5 years of endocrine therapy. A score below 18 indicates low risk of distant recurrence and claims to predict little to no effect of chemotherapy on patient outcomes. A score between 18 and 30 indicates intermediate risk of recurrence and claims to predict no substantial effect of chemotherapy on patient outcomes.
outcomes. A score of 31 or more indicates high risk of recurrence and claims to predict a large effect of chemotherapy on patient outcomes.

3.16 The breast recurrence score can be combined with clinical and pathological factors using the recurrence score-pathology-clinical (RSPC) calculator. However, this calculator has not been validated in a cohort independent of that used to derive Oncotype DX.

3.17 During consultation on the first diagnostics consultation document NICE accepted the company’s commitment to maintain the current confidential discount, which is in line with the Diagnostics Assessment Programme interim addendum on access proposals. This provides a simple discount to the list price of Oncotype DX, with the discount applied at the point of purchase or invoice. The level of the discount is commercial in confidence.

**Prosigna (NanoString Technologies)**

3.18 Prosigna is a CE-marked assay designed to provide information on breast cancer subtype and to predict distant recurrence-free survival at 10 years. The test is for postmenopausal people with early breast cancer that is ER-positive, HER2-negative and LN-negative or LN-positive (up to 3 positive nodes).

3.19 Prosigna measures the expression of 50 genes used for intrinsic subtype classification, 8 housekeeping genes used for signal normalisation, 6 positive controls, and 8 negative controls.

3.20 The test needs RNA extracted from a FFPE breast tumour tissue sample. It is based on direct mRNA counting using fluorescent probes and an nCounter Digital Analyser.

3.21 Prosigna classifies the risk of distant recurrence within 10 years, assuming 5 years of endocrine therapy, based on the PAM50 gene signature, breast cancer subtype, tumour size, nodal status and
proliferation score. The proliferation score is determined by evaluating multiple genes associated with the proliferation pathway. The test gives a score between 0 and 100. Based on this score and the nodal status, samples are classified into risk categories:

- LN-negative: low risk (0 to 40), intermediate risk (41 to 60) or high risk (61 to 100).
- LN-positive (up to 3 positive nodes): low risk (0 to 15), intermediate risk (16 to 40), or high risk (41 to 100).

3.22 During consultation on the first diagnostics consultation document, NICE accepted an access proposal from the company in line with the Diagnostics Assessment Programme interim addendum on access proposals. This provides a simple discount to the list price of Prosigna, with the discount applied at the point of purchase or invoice. The level of the discount is commercial in confidence.

**IHC4 and IHC4+C**

3.23 The IHC4 test is a laboratory developed test that combines the results of 4 immunohistochemistry (IHC) measurements. The IHC4+C test combines the results of the 4 IHC4 tests with clinical and pathological features such as age, nodal status, tumour size, and grade. Both versions are designed to quantify the 10-year risk of distant disease recurrence, assuming 5 years of endocrine therapy. The test is for postmenopausal people with early breast cancer that is ER-positive and LN-negative or LN-positive (up to 3 positive nodes).

3.24 The IHC4+C test needs an FFPE breast tumour tissue sample. The 4 immunohistochemistry tests are: ER, progesterone receptor (PR), HER2 and the proliferation marker Ki67. ER and HER2 markers are commonly measured in NHS laboratories, but PR and Ki67 markers are not.
3.25 The IHC4+C test is in clinical use at 1 NHS centre (the Royal Marsden NHS Foundation Trust), which carries out the test with an average turnaround time of 1 week. The test could be run in local NHS laboratories providing that training and quality assurance programmes for the individual assays are in place.

3.26 The IHC4+C test uses a published algorithm to calculate a risk score for distant recurrence based on the results of the 4 assays and clinical factors. A calculator is available for use on request. A score of less than 10% is categorised as low risk for distant recurrence at 10 years. A score of more than 10% but less than 20% is intermediate risk, and a score of 20% or more is high risk for distant recurrence at 10 years.

**The comparator**

3.27 The comparator is decision-making for adjuvant chemotherapy prescribing, based on clinical and pathological features or the results of tools used to assess risk without the tumour profiling tests. Features may include the stage of the disease, nodal status, ER or PR status, HER2 status and any previous treatment (for example, neoadjuvant therapy). Risk assessment tools include PREDICT, the Nottingham Prognostic Index (NPI) and Adjuvant! Online. However, Adjuvant! Online is currently unavailable because it is being updated. It is not certain when it will be reinstated, and the website directs people to the PREDICT tool.

3.28 These risk assessment tools can be used to define the level of clinical risk. For example, in LN-negative disease a NPI of 3.4 or less is classed as low risk, and a NPI of more than 3.4 is classed as intermediate risk. If using the PREDICT tool, an absolute 10-year survival benefit from chemotherapy of less than 3% is classed as low risk; between 3 and 5% is classed as intermediate risk; and more than 5% is classed as high risk.
## Evidence

The diagnostics advisory committee (section 8) considered evidence on EndoPredict, MammaPrint, Oncotype DX, Prosigna and IHC4 or IHC4+C from several sources. Full details of all the evidence are in the [committee papers](#).

### Clinical effectiveness

#### 4.1 Evidence on the following outcomes was of interest in the clinical effectiveness review:

- **Prognostic ability** – the degree to which the test can accurately predict the risk of an outcome such as disease recurrence.
- **Prediction of relative treatment effect** – the ability of the test to predict which patients have disease that will respond to chemotherapy. It can be assessed by considering whether the relative treatment effect of chemotherapy or no chemotherapy on patient outcomes differs according to the test score.
- **Clinical utility** – the ability of the prospective use of the test to affect patient outcomes such as recurrence and survival compared with current practice.
- **Decision impact** – how the test influences decision-making in terms of which patients will be offered chemotherapy.

#### 4.2 A total of 153 references were included in the review. Studies assessing prognostic ability and prediction of relative treatment effect were quality assessed using relevant criteria from the draft prediction model study risk of bias assessment tool (PROBAST). Clinical utility studies were quality assessed using the Cochrane risk of bias tool for randomised controlled trials (RCTs).

### Prognostic ability

#### 4.3 Studies providing information on prognostic ability were retrospective analyses of RCT data or routinely collected data. Most of the studies excluded patients who did not have a large...
enough tissue sample for testing, which leaves the evidence base at potential risk of spectrum bias, because patients with smaller tumours (who may be systematically different to those with large tumours) are likely to be under-represented. In many studies patients had chemotherapy, which could affect event rates and therefore potentially reduce the apparent prognostic performance of a test. In other studies, patients who had chemotherapy were excluded from analyses, which may also lead to spectrum bias. Therefore studies in which all patients had endocrine monotherapy were preferable.

4.4 Results for prognostic ability were generally presented as unadjusted or adjusted analyses. Unadjusted analyses look at differences in the event rates among low-, intermediate- and high-risk groups without adjusting for clinical and pathological variables. Adjusted analyses show whether the test has prognostic value over clinical and pathological variables.

**Distribution of patients across risk categories**

4.5 Among studies of patients with lymph node (LN)-negative disease who had endocrine monotherapy, in each group around 70% to 80% had disease that was categorised as low or low/intermediate risk across all tests (11 studies). Most MammaPrint studies had mixed endocrine and chemotherapy use, mixed hormone receptor status with or without mixed human epidermal growth factor receptor 2 (HER2) status, so results may not be comparable with results from other tests. In these studies 20% to 61% of patients had disease that was categorised as low risk (6 studies). Most IHC4 or IHC4+C studies used quartiles or tertiles to define risk groups. These studies do not provide useful information on the distribution of patients across risk categories.
4.6 The proportion of patients with low and intermediate risk was generally much lower in groups with LN-positive disease than in groups with LN-negative disease who had endocrine monotherapy (7 LN-positive studies). For Oncotype DX, however, the proportion of patients with low and intermediate risk was only slightly lower in the LN-negative group than in the LN-positive group. Studies of MammaPrint in patients with LN-positive disease were all done in groups with mixed hormone receptor status and mixed or unknown HER2 status, so results may not be comparable with results from other tests. In these studies 38% to 41% of patients had disease that was categorised as low risk (2 studies).

**Oncotype DX**

4.7 There were 11 data sets that provided information on the prognostic ability of Oncotype DX: 7 reanalyses of RCT data and 4 retrospective studies of routinely collected data. All studies were validation studies, and in 4 studies patients had endocrine monotherapy. Three of the studies were done in East Asia and may not be generalisable to England because usual clinical practice may differ between countries enough to affect prognostic outcomes. Also, it is possible that people of different ethnicities have different underlying risk profiles and natural history of disease.

4.8 Unadjusted analyses indicated that Oncotype DX had prognostic accuracy (there were statistically significant differences between low-risk and high-risk groups) across various recurrence outcomes, regardless of lymph node status. However, hazard ratios between the intermediate-risk group and the high- or low-risk groups were not always statistically significant, particularly in the group with LN-positive disease.

4.9 In adjusted analyses, Oncotype DX provided statistically significant additional prognostic information over most commonly used clinical
and pathological variables (age, grade, size, nodal status), regardless of lymph node status. A bespoke analysis of TransATAC study data also showed that Oncotype DX provided additional prognostic information over clinical and pathological tools to assess risk.

**MammaPrint**

4.10 There were 10 data sets that provided information on the prognostic ability of MammaPrint: 1 reanalysis of RCT data and 9 retrospective studies of routinely collected data. In addition, a further 4 studies pooled data on specific patients from the same 10 data sets. All studies were validation studies, and in 5 studies patients had endocrine monotherapy. Most studies included some patients who were out of scope (with HER2-positive or hormone receptor-negative disease or both).

4.11 In 6 of 7 unadjusted analyses, MammaPrint had prognostic accuracy (there were statistically significant differences between low-risk and high-risk groups) for 10 year distant recurrence-free survival or interval, regardless of LN status.

4.12 In adjusted analyses, a pooled analysis of patients with LN-negative and LN-positive disease showed that MammaPrint had statistically significant prognostic accuracy for 10-year distant recurrence-free survival after adjusting for clinical and pathological variables. In patients with LN-negative disease, MammaPrint had statistically significant prognostic accuracy for 10-year distant recurrence-free interval when adjusted for Adjuvant! Online or Nottingham Prognostic Index (NPI). In patients with LN-positive disease, MammaPrint had borderline statistically significant prognostic accuracy for 10-year distant recurrence-free survival when adjusted for clinical and pathological variables.
Prosigna

4.13 There were 8 data sets that provided information on the prognostic ability of Prosigna: 6 reanalyses of RCT data and 3 retrospective analyses of 2 prospective cohort studies. All studies were validation studies, and in 5 studies patients had endocrine monotherapy. Some studies included some patients who were out of scope (with HER2-positive or hormone receptor-negative disease or both).

4.14 Prosigna had statistically significant prognostic accuracy for 10-year distant recurrence-free survival and interval in all unadjusted analyses of patients with LN-negative and LN-positive disease.

4.15 In analyses adjusted for clinical and pathological variables or tools, Prosigna had prognostic accuracy for 10-year distant metastasis-free survival and distant recurrence-free survival. In patients with LN-negative disease the results were statistically significant. In patients with LN-positive disease the results were statistically or borderline significant.

EndoPredict

4.16 There were 3 data sets that provided information on the prognostic ability of EndoPredict; all were reanalyses of RCT data. All studies were validation studies, and in 2 of the 3 studies patients had endocrine monotherapy.

4.17 In unadjusted analyses, EndoPredict had statistically significant prognostic accuracy for 10-year distant recurrence-free survival and interval in patients with LN-negative and LN-positive disease.

4.18 Results from the bespoke analysis of TransATAC, which reported adjusted analyses on the EPclin score part of EndoPredict were academic in confidence. Two studies reported adjusted analyses on the EP score part of EndoPredict, showing that it provided
statistically significant additional information over clinical and pathological variables regardless of LN status.

**IHC4 and IHC4+C**

4.19 There were 12 data sets that provided information on the prognostic ability of IHC4 and IHC4+C: 6 reanalyses of RCT data and 6 reanalyses of routinely collected data. Most of the data related to the IHC4 score alone, without including clinical factors. One of the studies was based on the derivation cohort for IHC4, and therefore may have overestimated prognostic ability. The remaining studies were validation studies. Patients had endocrine monotherapy in only 2 studies, 1 of which was the derivation cohort study.

4.20 In unadjusted analyses, IHC4 had statistically significantly better prognostic performance in groups with high risk than in groups with low risk (defined by quartiles or tertiles) regardless of lymph node status. However, no studies reported survival or recurrence outcomes by risk group. Also, many used laboratory methods that differed from the derivation study methodology. In adjusted analyses, IHC4 had additional prognostic value over clinical and pathological factors in 3 studies, but patients had endocrine monotherapy in only 1 of these studies.

4.21 Data on IHC4+C came from the derivation cohort and 1 validation cohort. These studies showed that IHC4+C had prognostic value in unadjusted analyses. In adjusted analyses IHC4+C provided statistically significantly more information than the NPI in LN-negative, but not LN-positive, disease.

**Prediction of relative treatment effect**

4.22 In addition to estimating the risk of recurrence, the ability of Oncotype DX and MammaPrint to predict which patients have disease that will respond to chemotherapy was explored in studies.

Diagnostics consultation document: Tumour profiling tests to guide adjuvant chemotherapy decisions in early breast cancer

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The external assessment group (EAG) reviewed evidence in support of this.

**Oncotype DX**

4.23 In 5 data sets (2 reanalyses of RCT data and 3 observational studies) reported across 11 published references and 1 confidential manuscript, analyses assessed the ability of Oncotype DX to predict relative treatment effects for chemotherapy.

4.24 The 2 reanalyses of RCTs suggest that Oncotype DX may predict differences in relative treatment effects for chemotherapy. Hazard ratios for disease-free survival for patients having chemotherapy compared with those having no chemotherapy suggested that the greatest relative treatment effect was for patients in the Oncotype DX high-risk category. Unadjusted interaction tests between Oncotype DX risk group and relative treatment effects were mainly statistically significant, but adjusted interaction tests were not always statistically significant, particularly in the group with LN-positive disease.

4.25 Results from the 3 observational studies were mixed and at high risk from confounding. One reported a statistically significant interaction test but this was only adjusted for a limited number of factors. Two others reported hazard ratios for chemotherapy compared with no chemotherapy; 1 study in patients with intermediate risk, and another in patients with high risk. Both of these studies reported statistically non-significant results.

4.26 The recurrence score-pathology-clinical (RSPC) algorithm incorporates Oncotype DX plus age, tumour size and grade. There was a non-significant interaction test result between relative chemotherapy treatment effects and RSPC risk group.
**MammaPrint**

4.27 Two studies reported the ability of MammaPrint to predict the relative treatment effects for chemotherapy. In a pooled analysis including patients with LN-negative and LN-positive disease, the effect of chemotherapy compared with no chemotherapy was statistically significant in the MammaPrint high-risk group but not in the low-risk group in unadjusted and adjusted analyses. Further, the interaction test for chemotherapy treatment and risk group was non-significant. In a pooled analysis of patients with LN-positive disease, there was a non-significant interaction between chemotherapy treatment and risk group.

**Clinical utility**

4.28 There were no clinical utility data available for EndoPredict, Prosigna or IHC4+C.

**Oncotype DX**

4.29 Five data sets, reported across 9 published references and 1 confidential manuscript, reported evidence on the clinical utility of Oncotype DX. One further study did not meet the inclusion criteria (because of insufficient follow-up length), but presented subgroup data according to age, lymph node status and ethnicity, and was therefore discussed by the EAG. Studies generally reported different outcomes, making comparisons across studies difficult. The EAG noted that the best evidence for clinical utility is an RCT of treatment guided by the test compared with treatment guided by the comparator, and that this type of evidence is not currently available for Oncotype DX. All studies reporting on the clinical utility of Oncotype DX are judged to be of poor quality using the Cochrane risk of bias tool for RCTs.

4.30 In patients with LN-negative disease, using the test in clinical practice appeared to result in low rates of chemotherapy in patients
with low risk (2% to 12%), with acceptable outcomes (distant recurrence-free survival, distant recurrence-free interval or invasive disease-free survival 96% to 99.6%). Rates of chemotherapy increased with increasing risk category, and were generally higher in patients with LN-positive disease. It was not possible to conclude whether patients in intermediate and high-risk categories had better outcomes as a result of using Oncotype DX to guide treatment because there were no comparator groups (patients who had treatment without Oncotype DX testing).

### MammaPrint

4.31 Two studies reported evidence relating to the clinical utility of MammaPrint. MINDACT was a prospective, partially randomised study in which clinical risk was determined using a modified version of Adjuvant! Online. Patients with risk scores that disagreed from MammaPrint and modified Adjuvant! Online were randomised to chemotherapy or no chemotherapy. Of patients included in the study, 88% had HR-positive disease and 90% HER2-negative disease, therefore some patients were outside of the scope for this assessment. For the group who were high risk with modified Adjuvant! Online and low risk with MammaPrint, 5-year distant metastasis-free survival was 95.9% with chemotherapy and 94.4% without chemotherapy, a non-statistically significant absolute difference of 1.5% (adjusted hazard ratio for distant metastasis or death with chemotherapy compared with no chemotherapy, 0.78; 95% CI 0.50 to 1.21; p=0.27). For the group who were low risk with modified Adjuvant! Online and high risk with MammaPrint, 5-year distant metastasis-free survival was 95.8% with chemotherapy and 95.0% without chemotherapy, a non-statistically significant absolute difference of 0.8% (adjusted hazard ratio for distant metastasis or death with chemotherapy compared with no chemotherapy, 1.17; 95% CI 0.59 to 2.28; p=0.66). The EAG judged MINDACT to be at
4.32 Results from the RASTER study suggested that distant recurrence-free interval rates were sufficiently low in the MammaPrint low-risk group for these patients to avoid chemotherapy. The 5-year distant recurrence-free interval rate for LN-negative disease was 97.0% for patients with low risk (15% had chemotherapy) and 91.7% for patients with high risk (81% had chemotherapy). In addition, MammaPrint provided additional prognostic information over Adjuvant! Online and the NPI, but not over the NHS PREDICT tool. The EAG judged RASTER to be at high risk of bias using the Cochrane risk of bias tool for RCTs.

Comparison of the tests with each other

4.33 There were 6 studies that compared more than 1 test: 4 reanalyses of RCTs and 2 observational studies. Evidence shows that generally when a test placed more patients in a low-risk category than another test, the event-free survival in the low-risk group was reduced. Also, the tests generally performed differently in patients with LN-negative and LN-positive disease.

4.34 Thirteen studies reported data from microarray analyses on more than 1 test, however, these studies had methodological limitations. The comparability of test algorithms applied to microarray data with the commercial assays was unknown, so the generalisability of findings from microarray studies to the decision problem was uncertain. All the studies reported data on Oncotype DX and MammaPrint, and 2 also reported data on EndoPredict. The microarray studies generally supported the conclusions from studies using the commercial versions of the assays in suggesting that Oncotype DX, MammaPrint and EndoPredict can discriminate between patients with high and low risk regardless of LN status.
terms of additional prognostic performance of the tests over clinical and pathological variables, EndoPredict appeared to have the greatest benefit, followed by Oncotype DX and then MammaPrint. However, because of the methodological limitations, the EAG judged that these studies did not provide conclusive evidence of the superiority of 1 test over others.

4.35 The OPTIMA Prelim study, a UK-based feasibility phase of an RCT, analysed concordance between different tests. The study included Oncotype DX, MammaPrint, Prosigna and IHC4 plus 2 other tests. Out of the 4 in-scope tests, MammaPrint assigned the most patients to the low-risk category, but unlike the other 3 tests it does not have an intermediate category. When the low and intermediate categories were treated as 1 category for the 3 tests that have 3 risk groups, Oncotype DX assigned the most patients to this category, and MammaPrint the least. Kappa statistics indicated modest agreement between tests, ranging from 0.33 to 0.53. Also, across 5 tests in the study, only 39% of tumours were uniformly classified as either low/intermediate risk or high risk by all 5 tests. Of these, 31% were classified as low/intermediate risk by all tests and 8% were high risk by all tests. The study authors concluded that although the tests assigned similar proportions of patients to low/intermediate-risk and high-risk categories, test results for an individual patient could differ markedly depending on which test was used.

Decision impact

4.36 The review of decision impact focused on studies done in the UK or the rest of Europe:

- Oncotype DX: 6 UK studies and 12 other European studies
- EndoPredict: 1 UK study and 3 other European studies
- IHC4+C: 1 UK study and 0 other European studies
- Prosigna: 0 UK studies and 3 other European studies
- MammaPrint: 0 UK studies and 8 other European studies.

4.37 The percentage of patients with any change in treatment recommendation or decision (either to or from chemotherapy) in UK studies was 29% to 49% across 4 Oncotype DX studies, 37% in 1 EndoPredict study and 27% in 1 IHC4+C study. Ranges across European (non-UK) studies were 5% to 70% for Oncotype DX, 38% to 41% for EndoPredict, 14% to 41% for Prosigna and 13% to 51% for MammaPrint.

4.38 The net change in the percentage of patients with a chemotherapy recommendation or decision (pre-test to post-test) among UK studies was a reduction of 8% to 23% across 4 Oncotype DX studies, an increase of 1% in 1 EndoPredict study, and a reduction of between 2% and 26% in 1 IHC4+C study. Net changes across European (non-UK) studies were a reduction of 0% to 64% for Oncotype DX, a reduction of 13% to 26% for EndoPredict, a reduction of 2% to an increase of 9% for Prosigna, and a reduction of 31% to an increase of 8% for MammaPrint.

Anxiety and health-related quality of life

4.39 There were 6 studies that reported outcomes relating to anxiety (including worry and distress) and health-related quality of life. The lack of a comparator in the studies made it difficult to tell whether changes in anxiety experienced with the use of tumour profiling tests would also have occurred if patients received a definitive decision based on clinical risk factors alone. Overall, evidence suggests that tumour profile testing may reduce anxiety in some patients in some contexts, but generally there was little effect on health-related quality of life.
Cost effectiveness

Review of economic evidence

4.40 The EAG reviewed existing studies investigating the cost effectiveness of tumour profiling tests to guide treatment decisions in people with early breast cancer, and also did a detailed critique of the economic models and analyses provided by Agendia (MammaPrint), Genomic Health (Oncotype DX), and the chief investigator of a UK decision impact study (EndoPredict).

4.41 From the review, 26 studies were identified that had been published since the original assessment for diagnostics guidance 10. The models reported in the studies assessed the cost effectiveness of tumour profiling tests across different countries including the UK, the US, Canada, Mexico, Japan, Austria, Germany, France and the Netherlands. Most studies compared Oncotype DX (18 studies), MammaPrint (8 studies) or EndoPredict (1 study) with comparators such as Adjuvant! Online, the St Gallen guidelines, standard practice or other conventional diagnostic tools. There was variation between the analyses in the populations evaluated, the disease type and other patient characteristics.

4.42 There was a high level of consistency in the general modelling approach and structure, and several studies were based on a previously published model. Most of the models used a Markov or hybrid decision tree–Markov approach, 2 studies used a partitioned survival approach and 1 study used a discrete event simulation approach. The time horizons ranged from 10 years to the patient’s remaining lifetime, with cycle lengths ranging from 1 month to 1 year when reported. Most of the models that evaluated Oncotype DX assumed that the test could predict relative treatment effects for chemotherapy.
Economic evaluation

4.43 None of the models identified in the literature review included all of the tests identified in the scope. Therefore, the EAG developed a de novo economic model designed to assess the cost effectiveness of Oncotype DX, MammaPrint, Prosigna, IHC4+C and EndoPredict compared with current practice without the use of the tumour profiling tests. The model used a lifetime time horizon (42 years) from the perspective of the UK NHS and personal social services. All costs and health outcomes were discounted at a rate of 3.5% per year. Unit costs were valued at 2015/16 prices. The main source of evidence used to inform the analyses of Oncotype DX, Prosigna, IHC4+C and EndoPredict was a bespoke analysis of TransATAC provided by the study investigators. This was limited to UK data on patients with hormone receptor-positive, HER2-negative disease with 0 to 3 positive lymph nodes to match the scope for this assessment. Because this study did not include MammaPrint, MINDACT was used as the basis for evaluating the cost effectiveness of MammaPrint. PREDICT scores were not available in either data set, and so this tool could not be considered as a comparator or used to determine different risk subgroups. Therefore, the comparator for Oncotype DX, Prosigna, IHC4+C and EndoPredict was current practice (various tools and algorithms), and the comparator for MammaPrint was a modified version of Adjuvant! Online.

Model structure

4.44 The hybrid decision tree–Markov model was based on the model previously developed by Ward et al. (2013). The decision tree component of the model classified patients in the current practice group (no test) and the tumour profiling test group as high, intermediate and low risk. For EndoPredict and MammaPrint, the intermediate-risk category was excluded because the test provides
results in terms of high and low risk only. In both the test group and
the current practice group, the decision tree determined the
probability that a patient would be in 1 of 6 groups: low risk,
chemotherapy; low risk, no chemotherapy; intermediate risk,
chemotherapy; intermediate risk, no chemotherapy; high risk,
chemotherapy, and high risk, no chemotherapy. For EndoPredict
and MammaPrint, 4 groups were used because there was no
intermediate-risk category. Each group was linked to a Markov
model which predicted lifetime quality-adjusted life-years (QALYs)
and costs according to the patient’s risk of distant recurrence and
whether or not they had chemotherapy.

Each Markov node included 4 health states: distant recurrence-
free; distant recurrence; long-term adverse events (acute myeloid
leukaemia [AML]); and dead. Patients entered the model in the
distant recurrence-free health state. A health-related quality of life
decrement was applied during the first model cycle to account for
health losses associated with short-term adverse events for
patients having adjuvant chemotherapy. The treatment effect for
adjuvant chemotherapy was modelled using a relative risk
reduction for distant recurrence within each risk classification
group. The benefit of the test was therefore captured in the model
by changing the probability that patients with each test risk
classification had adjuvant chemotherapy.

Model inputs

The risk classification probabilities used in the model for Oncotype
DX, Prosigna, IHC4+C and EndoPredict were from the bespoke
data analysis of TransATAC, which only included postmenopausal
women. For MammaPrint, they were from MINDACT.

The probability of developing distant metastases in each group and
risk category was based on 10-year recurrence-free interval data
from the bespoke data analysis of TransATAC for Oncotype DX, Prosigna, IHC4+C and EndoPredict. For MammaPrint the probability of developing distant metastases was based on an adjusted analysis of 5-year distant metastasis-free survival data from MINDACT. The model assumed that the risk of distant metastases between 10 and 15 years was halved, and after 15 years was zero.

4.48 The probability of having chemotherapy in the current practice group and in the tumour profiling test groups was taken from the sources in table 1.
Table 1 Source for post-test probability of having chemotherapy

<table>
<thead>
<tr>
<th>Population</th>
<th>Source</th>
<th>Proportion of patients having chemotherapy</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Low risk</td>
</tr>
<tr>
<td><strong>Current practice group</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LN-negative, NPI≤3.4</td>
<td>NCRAS data set</td>
<td>0.07</td>
</tr>
<tr>
<td>LN-negative, NPI&gt;3.4</td>
<td>Genomic Health access scheme data set(^1)</td>
<td>0.43</td>
</tr>
<tr>
<td>LN-positive (1–3 nodes)</td>
<td>NCRAS data set</td>
<td>0.63</td>
</tr>
<tr>
<td>Overall population (MammaPrint)</td>
<td>Expert opinion</td>
<td>0.47</td>
</tr>
<tr>
<td><strong>3-level tests (Oncotype DX, Prosigna and IHC4+C)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LN-negative, NPI≤3.4</td>
<td>UKBCG survey data</td>
<td>0.00</td>
</tr>
<tr>
<td>LN-negative, NPI&gt;3.4</td>
<td>Genomic Health access scheme data set(^1)</td>
<td>0.01</td>
</tr>
<tr>
<td>LN-positive (1–3 nodes)</td>
<td>Loncaster et al. (2017) node-positive estimates</td>
<td>0.08</td>
</tr>
<tr>
<td><strong>2-level tests (EndoPredict and MammaPrint)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>EndoPredict: all 3 subgroups</td>
<td>Bloomfield et al. (2017) study</td>
<td>0.07</td>
</tr>
<tr>
<td>MammaPrint: all subgroups</td>
<td>Bloomfield et al. (2017) study</td>
<td>0.07</td>
</tr>
</tbody>
</table>

Abbreviations: LN, lymph node; NCRAS, National Cancer Registration and Analysis Service; NPI, Nottingham Prognostic Index; UKBCG, UK breast cancer group

\(^1\) The Genomic Health access scheme data set is based on the access scheme operated by NHS England and is a result of the research recommendation from NICE’s original diagnostics guidance 10

4.49 In the base-case analysis, the relative treatment effect for chemotherapy was assumed to be the same across all test risk groups, that is, all tests were assumed to be associated with prognostic benefit only. For Oncotype DX, Prosigna, IHC4+C and
EndoPredict a 10-year relative risk of distant recurrence was estimated as 0.76 for chemotherapy compared with no chemotherapy (Early breast cancer trialists’ collaborative group 2012), and was assumed to apply to the groups with LN-negative and LN-positive disease. For MammaPrint the 10-year relative risk of distant recurrence was estimated to be 0.77 (MINDACT) for chemotherapy compared with no chemotherapy.

In sensitivity analyses the effect of assuming that Oncotype DX could predict relative treatment effects for chemotherapy was explored, based on the B20 study by Paik et al. (2006) and the SWOG-8814 study by Albain et al. (2010). For the group with LN-negative disease, the 10-year relative risks of distant recurrence with chemotherapy compared with no chemotherapy were 1.31, 0.61 and 0.26 for the low-, intermediate- and high-risk categories respectively. For the group with LN-positive disease, the 10-year relative risks of relapse with chemotherapy compared with no chemotherapy were 1.02, 0.72 and 0.59 respectively. It is possible that the no-chemotherapy arm of B20 may have overestimated the difference in response rates between low- and high-risk patients, because this arm was the derivation set for Oncotype DX. Therefore, additional sensitivity analyses in the group with LN-negative disease explored the impact of varying the relative chemotherapy treatment effect between risk groups on the incremental cost-effectiveness ratios (ICERs). Hazard ratios were based on naive indirect comparisons of the chemotherapy arms from the B20 study and the no-chemotherapy arms from the B14 study (hazard ratios for treatment effects with chemotherapy compared with no chemotherapy were 0.64, 0.75 and 0.35 for the low-, intermediate- and high-risk categories respectively), and the chemotherapy arms of the B20 study and the no-chemotherapy arms of the TransATAC study (hazard ratios for treatment effects with chemotherapy compared with no chemotherapy were 0.86,
0.88 and 0.49 for the low-, intermediate- and high-risk categories respectively).

4.51 Survival following distant recurrence was based on a median of 40.1 months from Thomas et al. (2009). From this, the 6-month probability of death following distant recurrence was estimated to be 0.098, assuming a constant rate. The rate of death following distant metastases was assumed to be the same across the different subgroups and across each test risk group.

4.52 The model assumed that 10.5% of patients entering the distant recurrence health state had previously had local recurrence, based on de Bock et al. (2009). The 6-month probability of developing AML was estimated to be 0.00025, based on Wolff et al. (2015). Survival following the onset of AML was estimated to be approximately 8 months; assuming a constant event rate gave a 6-month probability of death following AML of 0.53. Additional sensitivity analyses explored the effect of including congestive heart failure (average net lifetime QALY loss of 0.0385 and average net lifetime cost saving of £2 from Hall et al. 2017, using an excess congestive heart failure risk relative to that of the general population), permanent hair loss (disutility of 0.04495 from Nafees et al. 2008 applied to 15% of all patients having chemotherapy) and peripheral neuropathy (disutility of 0.02 from Shiroiwa et al. 2009 applied to 12% of all patients having chemotherapy) in the model.

Costs

4.53 The costs of the tumour profiling tests were based on company prices (see table 2).
Table 2 Test prices

<table>
<thead>
<tr>
<th>Test</th>
<th>List price</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oncotype DX</td>
<td>£2,580</td>
<td>Tests carried out in Genomic Health laboratory in US. Cost includes sample handling and customer service. A commercial-in-confidence discounted test cost was used in the model.</td>
</tr>
<tr>
<td>Prosigna</td>
<td>£1,970</td>
<td>Based on doing the test in an NHS laboratory, which includes the laboratory costs (£240), the Prosigna kit (£1,650) and the nCounter system (£194,600) and is based on 2,500 samples per lifetime of the nCounter system. A commercial-in-confidence discounted test cost was used in a scenario analysis to account for the access proposal.</td>
</tr>
<tr>
<td>EndoPredict</td>
<td>£1,500</td>
<td>Tests carried out in Myriad's laboratory in Munich. A commercial-in-confidence discounted test cost was used in a scenario analysis to account for the access proposal.</td>
</tr>
<tr>
<td>IHC4</td>
<td>£203</td>
<td>The cost was based on 2014 prices. The total cost of the test (£198) was uplifted using the HCHS indices to current prices.</td>
</tr>
<tr>
<td>MammaPrint</td>
<td>£2,326</td>
<td>Converted from Euros to UK pounds sterling, assuming an exchange rate of 1 British pound to 1.15 Euros.</td>
</tr>
</tbody>
</table>

Abbreviations: HCHS, hospital and community health services

4.54  The costs associated with adjuvant chemotherapy were from a previous costing analysis of the OPTIMA Prelim trial (Hall et al. 2017). The weighted mean cost of adjuvant chemotherapy acquisition, delivery and toxicity was estimated to be £3,145 per course.

4.55  All surviving patients had endocrine therapy for a period of between 5 and 8 years. Costs of endocrine therapy were taken from the British national formulary (2017). In addition, 30% of women with early breast cancer had 4 mg of bisphosphonates (zoledronic acid) by intravenous infusion every 6 months for up to 3 years, at a cost of £58.50, excluding administration.
4.56 All patients had 2 routine follow-up visits during the first year after surgery, with annual visits thereafter for 5 years. Patients were also assumed to have a routine annual mammogram for up to 5 years. The cost of a routine follow-up visit was estimated to be £162.84, and the cost of a mammogram was estimated to be £46.37.

4.57 Costs associated with treating local recurrence were taken from Karnon et al. (2007) and uplifted to current prices (£13,913). This was applied as a once-only cost to distant recurrence. Costs associated with treating distant metastases were derived from Thomas et al. (2009), and included visits, drugs, pharmacy, hospital admission and intervention, imaging, radiotherapy, pathology and transport. Cost components specifically associated with terminal care were excluded. The 6-monthly cost of treating metastatic breast cancer was estimated to be £4,541.

Health-related quality of life

4.58 Health utilities were taken from published studies (see table 3).

Table 3 Health utilities applied in the base case

<table>
<thead>
<tr>
<th>Health state / event</th>
<th>Duration applied in model</th>
<th>Mean</th>
<th>Standard error</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Recurrence-free</td>
<td>Indefinite</td>
<td>0.824</td>
<td>0.002</td>
<td>Lidgren et al. 2007</td>
</tr>
<tr>
<td>Disutility distant metastases</td>
<td>Indefinite</td>
<td>0.14</td>
<td>0.11</td>
<td>Calculated from Lidgren et al. 2007</td>
</tr>
<tr>
<td>Local recurrence</td>
<td>Once-only QALY loss applied on transition to distant recurrence state</td>
<td>-0.108</td>
<td>0.04 (assumed)</td>
<td>Campbell et al. 2011</td>
</tr>
<tr>
<td>Chemistry AEs</td>
<td>6 months</td>
<td>-0.038</td>
<td>0.004</td>
<td>Campbell et al. 2011</td>
</tr>
<tr>
<td>AML</td>
<td>Indefinite</td>
<td>0.26</td>
<td>0.04 (assumed)</td>
<td>Younis et al. 2008</td>
</tr>
</tbody>
</table>

Abbreviations: AEs, adverse events; AML, acute myeloid leukaemia; QALY, quality-adjusted life year
Base-case results

4.59 The following key assumptions were applied in the base-case analysis:

- Clinicians interpreted each of the 3-level tests in the same way (for example, an Oncotype DX high-risk score would lead to the same chemotherapy decision as a Prosigna high-risk score).
- Clinicians interpreted each of the 2-level tests in the same way (for example, a MammaPrint high-risk score would lead to the same chemotherapy decision as an EndoPredict high-risk score).
- The treatment effect for adjuvant chemotherapy was the same across all risk score categories for all tests.
- The prognosis of patients with AML and the costs and QALYs accrued within the AML state were independent of whether they had previously developed distant metastases.
- A disutility associated with adjuvant chemotherapy was applied once during the first model cycle only (while the patient is taking the regimen).
- Costs associated with endocrine therapy, bisphosphonates, follow-up appointments and mammograms were assumed to differ according to time since model entry.
- The model assumed that people entered at an age of around 60 years.

4.60 In the subgroup with LN-negative disease and a NPI of 3.4 or less, compared with current practice, the probabilistic model gave ICERs of:

- £147,419 per QALY gained (EndoPredict)
- £122,725 per QALY gained (Oncotype DX)
- £91,028 per QALY gained (Prosigna)
- £2,654 per QALY gained (IHC4+C).
In the subgroup with LN-negative disease and a NPI of more than 3.4, compared with current practice, the probabilistic model gave ICERs of:

- £46,788 per QALY gained (EndoPredict)
- £26,058 per QALY gained (Prosigna)
- Oncotype DX was dominated by current practice (that is, it was more expensive and less effective)
- IHC4+C was dominant over current practice (that is, it was less expensive and more effective).

In the population with LN-positive disease, compared with current practice, the probabilistic model gave ICERs of:

- £28,731 per QALY gained (Prosigna)
- £21,458 per QALY gained (EndoPredict)
- Oncotype DX was dominated by current practice
- IHC4+C was dominant over current practice.

In the overall MINDACT population, MammaPrint compared with modified Adjuvant! Online had an ICER of £131,482 per QALY gained. In the modified Adjuvant! Online high-risk subgroup, MammaPrint was dominated by current practice, and in the modified Adjuvant! Online low-risk subgroup, MammaPrint compared with current practice had an ICER of £414,202 per QALY gained.

The risk classification probabilities and the probability of having chemotherapy were combined in the model to estimate chemotherapy use with and without tumour profiling. The modelled chemotherapy use in the base case is shown in table 4.
Table 4 Modelled chemotherapy use with and without tumour profiling

<table>
<thead>
<tr>
<th>Test, subgroup compared with current practice</th>
<th>Chemotherapy use</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Test</td>
<td>No test</td>
<td>Net change</td>
</tr>
<tr>
<td>Oncotype DX</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LN0 NPI ≤ 3.4</td>
<td>0.076</td>
<td>0.072</td>
<td>0.004</td>
</tr>
<tr>
<td>LN0 NPI &gt; 3.4</td>
<td>0.273</td>
<td>0.430</td>
<td>-0.157</td>
</tr>
<tr>
<td>LN+ (1–3 nodes)</td>
<td>0.337</td>
<td>0.627</td>
<td>-0.290</td>
</tr>
<tr>
<td>IHC4+C</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LN0 NPI ≤ 3.4</td>
<td>0.030</td>
<td>0.072</td>
<td>-0.042</td>
</tr>
<tr>
<td>LN0 NPI &gt; 3.4</td>
<td>0.355</td>
<td>0.430</td>
<td>-0.075</td>
</tr>
<tr>
<td>LN+ (1–3 nodes)</td>
<td>0.554</td>
<td>0.627</td>
<td>-0.073</td>
</tr>
<tr>
<td>Prosigna</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LN0 NPI ≤ 3.4</td>
<td>0.075</td>
<td>0.072</td>
<td>0.003</td>
</tr>
<tr>
<td>LN0 NPI &gt; 3.4</td>
<td>0.435</td>
<td>0.430</td>
<td>0.005</td>
</tr>
<tr>
<td>LN+ (1–3 nodes)</td>
<td>0.709</td>
<td>0.627</td>
<td>0.082</td>
</tr>
<tr>
<td>EndoPredict</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LN0 NPI ≤ 3.4</td>
<td>0.140</td>
<td>0.072</td>
<td>0.068</td>
</tr>
<tr>
<td>LN0 NPI &gt; 3.4</td>
<td>0.438</td>
<td>0.430</td>
<td>0.008</td>
</tr>
<tr>
<td>LN+ (1–3 nodes)</td>
<td>0.603</td>
<td>0.627</td>
<td>-0.024</td>
</tr>
<tr>
<td>MammaPrint</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MINDACT overall population</td>
<td>0.319</td>
<td>0.466</td>
<td>-0.148</td>
</tr>
<tr>
<td>mAOL high risk</td>
<td>0.445</td>
<td>0.772</td>
<td>-0.327</td>
</tr>
<tr>
<td>mAOL low risk</td>
<td>0.191</td>
<td>0.159</td>
<td>0.033</td>
</tr>
</tbody>
</table>

Abbreviations: LN0, lymph node negative; LN+, lymph node positive, mAOL, modified Adjuvant! Online; NPI, Nottingham Prognostic Index

**Probabilistic sensitivity analyses**

4.65 The cost-effectiveness planes from the probabilistic sensitivity analyses showed considerable uncertainty in the cost-effectiveness estimates.

4.66 In the subgroup with LN-negative disease and a NPI of 3.4 or less, the only test with a non-zero probability of producing more net benefit than current practice at maximum acceptable ICERs of £20,000 and £30,000 per QALY gained was IHC4+C.
4.67 In the subgroup with LN-negative disease and a NPI of more than 3.4, at a maximum acceptable ICER of £20,000 per QALY gained, IHC4+C had a probability of 0.69 of being cost effective compared with current practice. For all other tests, the probability that the test was cost effective compared with current practice at this threshold was 0.24 or less. In the same subgroup, at a maximum acceptable ICER of £30,000 per QALY gained, IHC4+C had a probability of 0.67 and Prosigna had a probability of 0.60 of being cost effective compared with current practice. Oncotype DX had a probability of 0.04 and EndoPredict had a probability of 0.26 of being cost effective compared with current practice.

4.68 In the subgroup with LN-positive disease, IHC4+C had probabilities of 0.95 and 0.94 of being cost effective compared with current practice at maximum acceptable ICERs of £20,000 and £30,000 per QALY gained respectively. In the same subgroup at the same maximum acceptable ICERs, the probability of EndoPredict producing more net benefit than current practice ranged from 0.44 to 0.73. For Prosigna the range was 0.24 to 0.55. In this subgroup Oncotype DX had very low probabilities of producing more net benefit than current practice at the same maximum acceptable ICERs (0.01 or lower).

4.69 In the overall MINDACT population and in the subgroups, the probability that MammaPrint would be cost effective compared with current practice at maximum acceptable ICERs of £20,000 and £30,000 per QALY gained was approximately zero.

**Deterministic sensitivity analyses**

4.70 The EAG did deterministic sensitivity analyses, testing a wide range of plausible values of key parameters.

4.71 Deterministic sensitivity analysis results for Oncotype DX compared with current practice were:
• Subgroup with LN-negative disease and a NPI of 3.4 or less: ICERs remained over £34,000 per QALY gained across all analyses.

• Subgroup with LN-negative disease and a NPI of more than 3.4: Oncotype DX was either dominated or had an ICER of more than £35,000 per QALY gained across almost all analyses. The only exception was when Oncotype DX was assumed to predict relative treatment effects for chemotherapy. In this analysis, Oncotype DX dominated current practice.

• Population with LN-positive disease: Oncotype DX remained dominated across most analyses. The exceptions were when Oncotype DX was assumed to predict relative treatment effects for chemotherapy (it was dominant), and when the cost of chemotherapy was doubled (£3,700 saved per QALY lost).

4.72 Deterministic sensitivity analysis results for IHC4+C compared with current practice were:

• Subgroup with LN-negative disease and a NPI of 3.4 or less: ICERs remained below £16,000 per QALY gained across all analyses, except when post-test chemotherapy probabilities were derived from Holt et al. (2011; £36,259 per QALY gained). Also, IHC4+C dominated current practice when the cost of chemotherapy was doubled.

• Subgroup with LN-negative disease and a NPI of more than 3.4: IHC4+C dominated current practice or had an ICER below £6,000 per QALY gained across all scenarios.

• Population with LN-positive disease: IHC4+C dominated current practice across all but 1 scenario. When the probability of having chemotherapy was based on the UK breast cancer group (UKBCG) survey the ICER was £1,929 per QALY gained.
4.73 Deterministic sensitivity analysis results for Prosigna compared with current practice were:

- Subgroup with LN-negative disease and a NPI of 3.4 or less: ICERs were greater than £71,000 per QALY gained across all analyses.
- Subgroup with LN-negative disease and a NPI of more than 3.4: ICERs were below £34,000 per QALY gained across all analyses.
- Population with LN-positive disease: ICERs were below £38,000 per QALY gained across all analyses.

4.74 Deterministic sensitivity analysis results for EndoPredict compared with current practice were:

- Subgroup with LN-negative disease and a NPI of 3.4 or less: ICERs remained greater than £91,000 per QALY gained across all analyses.
- Subgroup with LN-negative disease and a NPI of more than 3.4: ICERs remained greater than £30,000 per QALY gained across all but 2 of the analyses. Exceptions were when the UKBCG survey was used to inform the probability of having chemotherapy (£25,250 per QALY gained), and when Cusumano et al. (2014) was used to inform the probability of having chemotherapy based on the EndoPredict test result (£26,689 per QALY gained).
- Population with LN-positive disease: ICERs remained below £30,000 per QALY gained across all scenarios.

4.75 Deterministic sensitivity analysis results for MammaPrint compared with current practice were:

- Overall MINDACT population: ICERs were estimated to be greater than £76,000 per QALY gained across all scenarios.
• Modified Adjuvant! Online high-risk subgroup: MammaPrint was dominated by current practice across almost all scenarios.
• Modified Adjuvant! Online low-risk subgroup: ICERs were greater than £161,000 per QALY gained across all analyses.

5 Committee discussion

5.1 The committee discussed current practice for making adjuvant chemotherapy prescribing decisions. The clinical experts explained that NHS clinical practice has changed since NICE’s diagnostics guidance 10 was published in 2013. Also, the PREDICT tool is now used by many NHS trusts rather than the Nottingham Prognostic Index (NPI). Adjuvant! Online is not currently available. The committee also heard that Oncotype DX is currently used in NHS clinical practice and may be used for a broader group than the population defined in the original diagnostics guidance 10, that is, people with oestrogen receptor (ER)-positive, human epidermal growth factor receptor 2 (HER2)-negative and lymph node (LN)-negative early breast cancer who are assessed as being at intermediate risk using existing risk assessment tools.

5.2 The committee discussed the potential benefits of the tumour profiling tests for people with early breast cancer who are deciding whether to have adjuvant chemotherapy. It heard that there is potential benefit for people with cancer identified as being at low clinical risk, when test results suggest a high risk of distant recurrence. These people would therefore benefit from chemotherapy. It also heard that there is potential benefit for people with cancer categorised as high clinical risk, when test results suggest a low risk of distant recurrence. The committee heard that these people could decide not to have chemotherapy, therefore avoiding toxic side effects and effects on fertility. They could potentially resume normal daily activities earlier, although
some may wish to have chemotherapy regardless of the test result. However, the committee noted that the claimed benefits of the tests depend on them having sufficient accuracy and discrimination to correctly classify risk and provide valid clinical information. The clinical experts explained that the additional clinical information provided by the tests may help people discuss further treatment options. This information is particularly helpful for people with cancers identified as intermediate clinical risk when the decision to offer chemotherapy is unclear. However, the final decision to recommend a course of adjuvant chemotherapy would always take into account the person’s circumstances and preferences.

Clinical effectiveness

5.3 The committee considered the prognostic ability of the tumour profiling tests. It noted that for people with LN-negative disease, all the tests had statistically significant prognostic accuracy over clinical and pathological features or risk assessment tools such as the NPI (see section 4). It also noted that for people with LN-positive disease, results for prognostic ability were more variable but all tests except IHC4+C showed statistically significant or borderline statistically significant prognostic ability over clinical and pathological features or risk assessment tools. The external assessment group (EAG) explained that there were concerns about patient spectrum bias in all studies reporting prognostic ability. This was because in many of the studies some or all patients had chemotherapy or patients who had not had chemotherapy were selected for analyses. Also, most studies excluded tumour samples with insufficient tissue, and some studies included patients who had hormone receptor-negative or HER2-positive disease. The committee concluded that despite the potential spectrum bias, the evidence suggested that all the tumour profiling tests have the ability to predict the risk of distant recurrence in the population.
included in the assessment. It also concluded that the evidence was weaker in the group with LN-positive disease than in the group with LN-negative disease.

5.4 The committee considered the evidence on whether the tumour profiling tests can predict relative treatment effects associated with chemotherapy. The clinical experts stated that it is likely some patients could have a greater relative treatment effect from chemotherapy than others, for example, patients with hormone receptor-positive cancer that is not sensitive to endocrine therapy, but evidence is not available to support this. The EAG explained that the only evidence available to show a relative treatment effect for chemotherapy across different risk groups is for Oncotype DX, and this evidence is weak because it is at high risk of bias from potential confounding. The results of interaction tests (which show whether the tumour profiling test is able to predict a different treatment effect by risk group) in the adjusted analysis in the B20 study by Paik et al. (2006; LN-negative disease) remained statistically significant when adjusting simultaneously for clinical and pathological variables. However, the EAG also explained that the difference in relative treatment effects for chemotherapy in the B20 study may be overestimated because this was the Oncotype DX derivation data set. In the SWOG-8814 study by Albain et al. (2010; LN-positive disease) the results of the interaction tests remained statistically significant when adjusting for some individual clinical and pathological variables, but there was no analysis that adjusted for these simultaneously, and the test was non-significant when adjusting for Allred-quantified ER status. The clinical experts explained that hormone receptor status may also predict relative treatment effects for chemotherapy. The committee considered that if all known clinical and pathological variables were included in the analyses of the SWOG-8814 data then it was likely that the results of the interaction test would no longer be statistically significant.
This suggested highly uncertain relative treatment effects for chemotherapy according to the results of the tumour profiling tests for this group with LN-positive disease. The committee concluded that the evidence on the extent to which tumour profiling tests are able to predict relative treatment effects for chemotherapy is highly uncertain, but there may be some differences between Oncotype DX risk groups. The committee noted that no data were available to assess a difference in relative treatment effects for chemotherapy for EndoPredict, IHC4+C and Prosigna risk groups, and that data on MammaPrint suggest no difference in relative treatment effects for chemotherapy.

5.5 The committee considered the evidence on clinical utility, that is, data from studies which assessed the ability of the tumour profiling tests to affect patient outcomes. It noted that the only test with evidence from randomised controlled trials in which patients were randomised to treatment guided by either test result or usual clinical practice was MammaPrint. The committee noted that MINDACT was a well-designed study. The results suggested that patients with high clinical risk and MammaPrint low-risk scores can forgo chemotherapy without a statistically significant increase in the 5-year risk of distant recurrence. However, a clinical expert explained that the risk of recurrence often continues beyond 5 years and noted that the MINDACT authors (Cardoso et al. 2016) stated that long-term follow-up and outcome data will be essential. These data are being collected and a 10-year follow-up analysis is planned. The committee noted that none of the other tumour profiling tests had similar evidence of clinical utility, but it was aware that this evidence was being collected for Oncotype DX and Prosigna (see section 5.22). The committee concluded that none of the tests had strong enough evidence to demonstrate an effect on subsequent patient outcomes.
5.6 The committee was encouraged by the availability of the data set provided in confidence to NICE by Genomic Health. The data set was based on the access scheme operated by NHS England, which provided real world evidence on the use of adjuvant chemotherapy in the NHS following testing with Oncotype DX for the population included in the scope for this assessment. The committee noted that the total number of patients in the data set appeared to be much larger than the number of patients with complete data in the population of interest, and that the advice from clinical experts (see section 5.1) was that the test has been used on a wider group of patients in practice. The committee concluded that the access scheme data set was an important piece of real world evidence for use in the economic model, but that more complete data could have been collected and reported. It also concluded that future data collection should be done as part of a national database, rather than by individual companies, to increase transparency and link to outcome data (see section 5.24).

5.7 The committee discussed the analytical validity of IHC4+C. The EAG explained that the evidence has developed since diagnostics guidance 10 was published. The committee noted that the data showed good correlation between different centres when scoring and staining were assessed separately for measurement of the Ki67 marker, which had been achieved with training. But it also noted that when studies looked at staining and scoring combined, the correlation between centres decreased substantially. A clinical expert noted that different antibody clones are available for testing Ki67, ER and progesterone receptor (PR) status, and that different studies used different antibody clones which means that the studies are not directly comparable. The committee heard that different methods of assessing ER and PR receptors may be needed for IHC4+C compared with those already used routinely, which may introduce additional complexity. The committee concluded that
because of these issues, the reproducibility of IHC4+C is poor. It also concluded that if this test were to be developed further the antibody clones used in the assays for ER, PR and Ki67 should be specified, and there would need to be substantial investment in staff training and quality assurance.

**Cost effectiveness**

5.8 The committee discussed the assumptions and inputs used in the model, and carefully considered the extensive stakeholder comments on the model and EAG responses to these comments. It noted that a specific analysis of the TransATAC data was used for risk classification probabilities and for distant recurrence rates based on test result for Oncotype DX, EndoPredict, Prosigna and IHC4+C. The results from this specific analysis of the data set have now been published. The EAG explained that this data source was chosen because it included data on 4 of the 5 tests of interest and was specific to the population included in the scope (patients with hormone receptor-positive, HER2-negative disease). The committee heard that although the TransATAC data were slightly older and some patients were not candidates for chemotherapy, the patient characteristics matched well with the more recent MINDACT study. The alternative would be to use different data sources for each test, which would have introduced additional uncertainty and complexity. Also, the group with LN-negative disease could not have been split according to level of clinical risk. The EAG also explained that the distant recurrence-free rates from the TransATAC analysis used in the model were consistent with results from other studies (B14, B20, TAILORx, MD Anderson, Clalit, Memorial Sloan Kettering, SEER and WSG PlanB) both when grouped separately by clinical risk and when all clinical risk groups were pooled together. The committee concluded that the
TransATAC analysis had some limitations, but was the best available data for use in the model.

5.9 The committee considered the data on pre- and post-test chemotherapy decisions used in the model. The EAG explained that for 3-level tests (tests with low, intermediate and high-risk categories [IHC4+C, Oncotype DX, Prosigna]), data on pre- and post-test chemotherapy decisions for the group with LN-negative disease and a NPI of more than 3.4 were taken from the Genomic Health access scheme data set (see section 5.6). For other clinical risk subgroups with the 3-level tests, and for all clinical risk subgroups with 2-level tests (tests with low and high-risk categories [EndoPredict, MammaPrint]), data on pre-test chemotherapy decisions were taken from different sources to data on post-test chemotherapy decisions. There were also very limited UK data for these groups. The committee considered the modelled impact of these data on chemotherapy use, and noted that although clinical and patient experts thought that the main benefit of the tests was in avoiding unnecessary chemotherapy, most tests were estimated to increase chemotherapy use at least in some subgroups (see section 4.48). The committee concluded that there was much more uncertainty around chemotherapy decision-making for the 2-level tests, and for the subgroups who were not included in the original NICE recommendation on tumour profiling tests (LN-negative disease and a NPI of 3.4 or less, and LN-positive disease).

5.10 The committee considered how adjuvant chemotherapy treatment effects had been applied in the economic model, particularly the relative treatment effects of chemotherapy between the risk groups predicted by the tumour profiling tests. It noted its earlier conclusion that the evidence on whether tumour profiling tests can predict relative treatment effects for chemotherapy is highly uncertain, but that there may be some differences between Oncotype DX risk
groups (see section 5.4). It agreed that for EndoPredict, IHC4+C and Prosigna, no evidence was available to show a difference in relative treatment effects of chemotherapy across risk groups, and that data on MammaPrint suggested no difference in relative treatment effects. Therefore for these tests it was appropriate to assume the same relative risk of distant recurrence across all test risk categories (0.76). The committee considered stakeholder comments submitted during the first consultation suggesting that Oncotype DX has the ability to predict which patients have disease that will respond to chemotherapy. The EAG noted that in response to the comments it had done additional exploratory analyses for Oncotype DX to show the impact on the incremental cost-effectiveness ratios (ICERs) if a smaller relative treatment effect than that taken from the B20 study (Paik et al. 2006) was applied in the model in the group with LN-negative disease and a NPI of more than 3.4 (see section 4.50). The EAG noted that the hazard ratios used in these analyses were from comparisons of independent arms of trials and were therefore very uncertain. The EAG also said that using hazard ratios calculated from the B20 (Paik et al. 2006) and the B14 (Paik et al. 2004) studies resulted in an ICER of around £24,000 per quality-adjusted life year (QALY) gained for Oncotype DX compared with current practice. Using hazard ratios calculated from the B20 and TransATAC studies resulted in an ICER of around £8,000 per QALY gained. The committee concluded that although these analyses were associated with considerable uncertainty, they gave an indication of Oncotype DX’s likely cost effectiveness if the relative treatment effects for chemotherapy did differ between Oncotype DX risk groups, but not to the extent reported in the Paik et al. (2006) study.

5.11 The committee considered stakeholder comments submitted during the first consultation suggesting that adverse events had not been adequately captured in the economic model; in particular,
congestive heart failure, permanent hair loss and peripheral neuropathy. The EAG noted that in response to the comments it had done additional exploratory analyses to include these adverse events in the model. Congestive heart failure was added into the model by incorporating estimated lifetime QALY losses and costs taken from an alternative model (Hall et al. 2017). Hair loss and peripheral neuropathy were incorporated using a disutility applied to a proportion of the population for the lifetime of the model. The EAG highlighted the considerable limitations of these analyses, and noted that for tests that increased chemotherapy use in some subgroups, the ICERs became less favourable. The committee noted that including additional adverse events in the model did reduce some of the ICERs, but not enough to change the conclusions. It also noted a further EAG analysis, which suggested that for tests that reduced chemotherapy use but were not cost effective, the QALY gain from avoiding adverse events would have to be in the range of 1.1 to 1.3 to result in cost-effective ICERs. The committee concluded that it was important to consider potential adverse events that could be caused by chemotherapy. However, the reduction in adverse events from reduced chemotherapy use, while beneficial for patients, was unlikely to affect its conclusions on the cost effectiveness of the tumour profiling tests based on the EAG’s analysis.

5.12 The committee considered other assumptions used in the model such as the cost of chemotherapy, how the risk of distant recurrence was applied over time, and adjuvant chemotherapy treatment effects if a different relative risk across risk groups is not assumed. The EAG explained that there was some uncertainty around these inputs, but all had been tested in sensitivity analyses. The committee concluded that the assumptions and inputs used in the model were reasonable, but they were associated with
considerable uncertainty because of the limitations in the data that underpinned them.

5.13 The committee noted its discussion on current practice (see section 5.1) and considered the absence of comparisons of the tumour profiling tests with the PREDICT tool. The EAG explained that in the model it was not possible to compare the tumour profiling tests with PREDICT, or to define the clinical risk groups using PREDICT, because relevant data were not available. The committee noted that the comparisons in the model do not fully reflect current NHS clinical practice, which leads to uncaptured uncertainty in the model results. The committee concluded that research on tumour profiling tests should include comparisons with PREDICT (see section 5.22) so that the cost effectiveness of the tests relative to current practice can be fully assessed in future.

5.14 The committee considered the subgroups that were included in the model, that is, people with LN-negative disease and a NPI of 3.4 or less, LN-negative disease and a NPI of more then 3.4, and LN-positive disease. It noted its earlier conclusion that the evidence suggested that all the tumour profiling tests have the ability to predict risk of distant recurrence (prognosis), but this ability was less certain in the group with LN-positive disease (see section 5.3). The committee also recalled that the test results are particularly helpful for people with cancers identified as intermediate clinical risk when the decision to offer chemotherapy is unclear (see section 5.2). The clinical experts explained that tumour profiling tests are also helpful for people with LN-positive cancer, who have comorbidities and therefore an additional reason to want to avoid chemotherapy. The EAG noted that this subgroup of the LN-positive population could not be modelled because of a lack of data. In addition, the committee noted that the EAG’s systematic review had highlighted substantial lack of agreement between the
tests in risk categorising the group with LN-positive disease. The committee decided to consider the ICERs in the group with LN-negative disease only, but noted that further studies would be helpful to assess the clinical effectiveness of the tests in the group with LN-positive disease (see section 5.23).

5.15 The committee considered the results from the model. It noted that the differences in the QALYs were small, and that the ICERs for all tumour profiling tests were highly uncertain because of the available clinical data and the assumptions used in the modelling (see sections 5.8 to 5.12). It also noted that the base-case ICERs for many of the tumour profiling tests were higher than those normally considered to be cost effective. However, it heard that access proposals had been made by Myriad Genetics (for EndoPredict) and NanoString Technologies (for Prosigna). Genomic Health confirmed that the confidential discount for Oncotype DX would continue in the NHS. The committee concluded that the availability of the access proposals for EndoPredict and Prosigna may reduce the ICERs to a range that could be considered plausibly cost effective despite the clinical uncertainties.

5.16 The committee considered the EndoPredict and Prosigna access proposals. Compared with current practice, the ICERs for EndoPredict and Prosigna in the group with LN-negative disease and a NPI of 3.4 or less were still higher than those normally considered to be a cost-effective use of NHS resources. In the group with LN-negative disease and a NPI of more than 3.4, Prosigna compared with current practice had an ICER of less than £20,000 per QALY gained, and therefore could be considered cost effective. In the same group, EndoPredict compared with current practice had ICERs between £20,000 and £30,000 per QALY gained, and varied depending on whether the testing was done at a
local or a centralised laboratory. The committee noted that localised testing was more cost effective than centralised testing, and that testing became more cost effective as test throughput increased. It also recalled its conclusion that the data on post-chemotherapy decisions were more uncertain for 2-level tests than for 3-level tests (see section 5.9), and noted that the EAG’s sensitivity analyses using plausible alternative sources for post-chemotherapy decisions resulted in ICERs that were lower than £20,000 per QALY gained. The committee decided that although there is uncertainty around the ICERs for EndoPredict compared with current practice, sensitivity analyses suggested that the ICER will be around £20,000 per QALY gained, and therefore it could be considered cost effective. The committee concluded that EndoPredict and Prosigna, when provided at the costs stated in the access proposals, were likely to be cost effective in the group with LN-negative disease and a NPI of more than 3.4, but evidence on clinical outcomes will be important to confirm this (see section 5.24).

5.17 The committee considered the ICERs for Oncotype DX compared with current practice. It heard that the proposed confidential test cost for Oncotype DX was the same as in current NHS practice, and that this cost had been used in the EAG’s economic model. It noted that in the base-case analyses Oncotype DX was dominated by the comparator in the group with LN-negative disease and a NPI of more than 3.4. The committee recalled its earlier conclusions; Oncotype DX may be able to predict relative treatment effects for chemotherapy, and the ICERs for Oncotype DX compared with current practice when some relative treatment effect across different risk groups was applied in the model were most likely to be between £9,000 and £25,000 per QALY gained (see section 5.4). However, it noted that this was very uncertain. The committee concluded that Oncotype DX, when provided at the test
cost stated in the access proposal, was likely to be cost effective in
the group with LN-negative disease and a NPI of more than 3.4, but
evidence on clinical outcomes will be important to confirm this (see
section 5.24).

5.18 The committee considered the ICERs for MammaPrint compared
with modified Adjuvant! Online. It noted that in the base-case
analyses, MammaPrint was dominated by the comparator in the
modified Adjuvant! Online high-risk subgroup. In the modified
Adjuvant! Online low-risk subgroup, the ICERs were much higher
than those normally considered to be cost effective. The committee
concluded that MammaPrint would not be a cost-effective use of
NHS resources.

5.19 The committee considered the ICERs for IHC4+C compared with
current practice. It noted that the ICERs were low or that IHC4+C
dominated current practice in all subgroups. The committee felt that
the test cost had been underestimated because it did not include
any costs for training or for setting up a quality assurance
programme. But even if these costs were included, IHC4+C may
still be cost effective. However, the committee noted its earlier
conclusion on the analytical validity of IHC4+C (see section 5.7)
and concluded that it could not be recommended for use in the
NHS until issues around reproducibility and implementation have
been resolved.

5.20 The committee noted that the model for EndoPredict, IHC4+C,
Oncotype DX and Prosigna related only to a postmenopausal
population because TransATAC was used as the data source for
these tests. It considered whether the model results could also
apply to a premenopausal population. A clinical expert explained
that the biology of a cancer and its molecular subtype, for example
hormone receptor status and HER2 status, is more influential in
determining the risk of distant recurrence than menopausal status. Therefore the committee concluded that the model results apply to premenopausal and postmenopausal populations.

5.21 The committee discussed the generalisability of the data to men. It acknowledged that men make up a small proportion of people with breast cancer. The committee noted that all the clinical and economic evidence was based on trials with women, but that the general subtypes are identical in men and women, and in clinical practice men would have treatment in the same way as women. The committee concluded that the recommendations in this guidance should also apply to men.

Research considerations

5.22 The committee noted that there are several ongoing studies which will provide evidence of long-term patient outcomes: further data collection from the MINDACT study on MammaPrint, the TAILORx trial on Oncotype DX and the OPTIMA trial on Prosigna. The committee concluded that these studies are relevant to this assessment and data from them may be important when the guidance is considered for updating in the future. But it noted that not all studies would provide UK-specific data and comparisons with the PREDICT tool, which would be important for future updates to fully assess the cost effectiveness of the tests compared with current practice.

5.23 The committee recalled its previous conclusion on the potential utility of the tests in the group with LN-positive disease (see section 5.14), particularly for people who have comorbidities and who may be particularly affected by the side effects of adjuvant chemotherapy. It noted that further research in this group would be welcome and heard from clinical experts that the ongoing OPTIMA
trial may help to reduce some of the uncertainties identified during this assessment.

**Data collection arrangements**

5.24 The recommendations for EndoPredict, Oncotype DX and Prosigna are conditional on data collection agreements being put in place. NICE will be inviting Genomic Health, Myriad Genetics and NanoString Technologies to review the proposed data collection agreements during the consultation period for this draft guidance. It is anticipated that companies will be asked to make arrangements to collect timely and complete record-level test data, which can be submitted to the National Cancer Registration and Analysis Service, with the aim of linking test data to chemotherapy use, recurrence and survival outcomes.

6 **Implementation**

NICE intends to develop tools, in association with relevant stakeholders, to help organisations put this guidance into practice.

7 **Review**

NICE reviews the evidence 3 years after publication to ensure that any relevant new evidence is identified. However, NICE may review and update the guidance at any time if significant new evidence becomes available.

Mark Kroese
Chair, diagnostics advisory committee
April 2018
8 Diagnostics advisory committee members and NICE project team

Diagnostics advisory committee

The diagnostics advisory committee is an independent committee consisting of 22 standing members and additional specialist members. A list of the committee members who participated in this assessment appears below.

Standing committee members

Dr Mark Kroese
Chair, diagnostics advisory committee

Mr John Bagshaw
In-vitro Diagnostics Consultant

Professor Enitan Carrol
Chair in Paediatric Infection, University of Liverpool

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Dr Steve Edwards
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NICE project team

Each diagnostics assessment is assigned to a team consisting of a technical analyst (who acts as the topic lead), a technical adviser and a project manager.