BREAST PATHOLOGY

An overview of assessment of prognostic and predictive factors in breast cancer needle core biopsy specimens

E A Rakha, I O Ellis

Needle core biopsy (NCB), as part of triple assessment for preoperative evaluation and diagnosis of breast cancer, is now considered as an established, highly accurate method for diagnosing breast cancer that has replaced either fine needle aspiration cytology or excisional biopsy as the initial diagnostic biopsy procedures in many institutions. In addition to its primary role in establishing an accurate histological diagnosis, NCB can potentially provide important additional pathological prognostic information which may be of direct clinical value in certain situations, such as patients being considered for preoperative (neoadjuvant) therapy. With this background in mind we briefly review the current role of NCB in breast cancer diagnosis and then concentrate this review on the usefulness and issues relating to use of this technique in providing accurate, reliable and clinically relevant preoperative prognostic and predictive information in patients with breast cancer.

Since Bolmgren et al\(^1\) introduced stereotactic needle biopsy of the breast in the late 1970s, and the introduction of automated core biopsy guns in the mid 1990s, needle core biopsy (NCB) has become an increasingly important diagnostic tool in the assessment of both palpable and non-palpable breast lesions.\(^2,3\) NCB is now considered the method of choice for tissue sampling as part of the triple assessment of breast disease,\(^3\) and the published data suggest that the use of core biopsy has increased the preoperative diagnosis rate in screen detected breast cancers.\(^3,4\) NCB, as compared to preoperative fine needle aspiration biopsy, can reliably distinguish between in-situ and invasive cancers, allow evaluation of more histological, prognostic, and predictive factors in breast cancer, and provide ample tissue for ancillary testing such as immunohistochemical studies (IHC), DNA and RNA analysis and other molecular techniques. In some patients with breast cancer, for example those treated with preoperative chemotherapy, the NCB specimen might be the only pretreatment tissue sample available for studies of prognostic and predictive markers. Moreover, NCB specimens can be evaluated easily by histopathologists to whom H&E stained sections are familiar.\(^7\)

NCB can be performed either freehand or with image guidance. Initial use of image guided NCB was reported with the utilisation of stereotactic imaging and extended to ultrasound-guided biopsies that facilitated biopsy of the clinically occult lesions. These biopsy specimens can be collected with automated (spring-activated) needle core or vacuum assisted biopsy device. However, the biopsy method used depends on which modality best depicts the abnormality, location and size of the abnormality within the breast, patient factors, and the operator’s experience and preference.

DIAGNOSTIC ACCURACY OF NCB

The diagnostic accuracy of NCB has been intensely verified and several studies have shown good concordance between NCB and subsequent surgical excision biopsy for diagnosis of carcinoma (ranging from 91% to 100%).\(^3,4\) The sensitivity for detection of malignancy is high in the majority of published studies (85–100%) and specificity is 96–100% for stereotactically guided NCB,\(^4,14–16\) although a few studies have shown a lower rate of sensitivity (eg 71%) and specificity (eg 85%). Slightly better results were documented in ultrasound-guided NCB and vacuum-assisted biopsies than in stereotactic, spring-loaded and freehand biopsies,\(^12,22–28\) and for diagnosis of palpable masses than for calcifications.\(^2\) However, the accuracy of NCB in the diagnosis of malignancy and the amount of information gained from it may be affected by several other factors, including basic definitions and methodology, case selection (palpable versus screen-detected lesion),\(^2,19\) needle size,\(^23–26\) number of cores taken,\(^19,20,23–26\) amount of clinical material obtained,\(^19\) presence of calcification on NCB,\(^33\) level of expertise of the operator and breast pathologist,\(^19,19,16\) and even timing of the biopsy.\(^31\) For example, some authors have reported that one of the major problems with assessing pathological factors on NCB specimens is undersampling of the most informative areas; therefore, an adequate volume of biopsy material is recommended, particularly during sampling of tumours with large size and in cases of calcifications.\(^22,41\) However, there are no available guidelines that indicate the number of cores or volume of tissues that should be obtained from tumours of certain size or with a specific amount of calcification. This should ideally be determined by common sense, and patient tailored. In our own clinic setting audit of ductal carcinoma in-situ (DCIS), diagnosis in patients presenting with mammographic calcification has shown that an optimum of either three or more cores containing calcium or five or more flecks of calcium in total is required to ensure adequate sampling of a calcification lesion.\(^42\) It is important that radiologists sampling breast lesions understand that both quality and...
quantity of tissue sampled are important to achieve high levels of sensitivity and specificity in needle core biopsy non-operative diagnosis.

**EVALUATION OF PROGNOSTIC AND PREDICTIVE FACTORS OF BREAST CARCINOMA ON NCB**

NCB is thus considered to be a highly accurate method of obtaining a preoperative histological diagnosis of breast cancer. There is dispute, however, about the range of prognostic and predictive information that should be obtained from NCB and included in the pathology report. Pathological prognostic and predictive factors are used in clinical practice for a variety of reasons. They provide detailed information on the prediction of outcome and response to therapy of an individual patient as a basis for accurate pretreatment planning of preoperative (neoadjuvant), operative and postoperative (adjuvant) therapy.

There are still insufficient data available at present to allow a consensus on what are reasonable performance goals for core biopsy diagnosis. However, it has been reported that once the diagnosis of malignancy is made on NCB, and when there is sufficient tissue, the following pathological features could be clearly identified in the pathology report. For invasive cancer: tumour type, histological grade, the presence, or absence of tumour necrosis. For DCIS: the extent of the in-situ proliferation, nuclear grade.

**HISTOLOGICAL PROGNOSTIC FACTORS**

**Tumour type**

Tumour type has been shown to be a prognostic factor in breast cancer. Histological typing of invasive breast tumours is carried out on well fixed samples and involves the assessment of several blocks to determine all of the features present. The evaluation of a variety of features including the extent of tubule formation, the presence of lymphoid stroma, mucin production, syncyrtial or discohesive growth pattern, and “Indian file” formation allows tumours to be separated according to type.

NCB can accurately predict histological type in the great majority of cases, with a rate of concordance with the full surgical specimen ranging from 93% to 100%. However, lower figures have also been reported (table 1).

**Tumour grade**

Histological grade is an important and well-established prognostic factor in breast cancer. Although different grading systems have been used to assess tumour grade, it has been any other available ancillary tests that can be performed on formalin-fixed paraffin embedded tissue.

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ER, oestrogen receptor; PgR, progesterone receptor.
reported that Nottingham modification of the Bloom and Richardson method, in which tubule formation, nuclear pleomorphism, and mitotic frequency are evaluated, gives reproducible results and correlates well with survival. Current evidence shows that grading on core biopsy can be performed and appears straightforward because formalin fixation is likely to be optimum. Concordance between grade on core biopsy and that in the definitive excision specimen can be achieved in approximately 75% of cases, varying in different studies from 59% to 91%. In a previous study (table 1), with the highest level of agreement achieved in grade 3 carcinomas.

Most of the discrepancy was that the surgical materials tended to be higher grade (upgrade) than NCB, almost invariably by only one level. Therefore, care should be taken with interpretation of grade on NCB. There may be insufficient amount of tumour in the core to allow 10 high power fields to be counted for mitosis; in surgical specimens, mitotic figures are counted in the periphery of the tumour, the growing edge of the tumour, where they are more frequent; there is evidence that the observed mitotic count may be lower in the core biopsy than in the excision specimen, resulting in a systematic bias leading to underscoring overall grade on NCB. In our own experience this latter issue is the main cause of discrepancy between NCB grade and final grade derived from examination of the surgical specimen. It is most probably due to traumatic damage of cells in the fragile state of mitosis obscuring their visibility in H&E stained sections. To date no effective routinely applicable alternatives have emerged for identification or scoring of mitosis frequency or growth fraction in core biopsy samples.

**Tumour size**

Tumour size is another well-established time dependent prognostic factor in breast cancer. In a comparison study between NCB and subsequent excision specimens from 79 women with invasive breast cancer, Sharifi et al. found a significant correlation between tumour size on NCB and excision specimens (r² = 0.30, p = 0.01); however, the pathological T stage was underestimated on NCB in 79% of cases and the T stage was underestimated on NCB in 71% of T1 lesions. Although grade on NCB, an approximate evaluation of the minimum size of the tumour may also contribute, measurement of this parameter on NCB is not recommended as a routine practice. If a preoperative estimate of size is required, this should be performed ultrasonographically.

**IN-SITU COMPONENT**

Previous studies of the accuracy of NCB have addressed the ability of NCB to detect in-situ disease in both non-palpable mammographically detected lesions and in palpable tumours. These studies have shown that the accuracy was lower for diagnosis of DCIS than for invasive disease. In addition, a serious problem may arise when a diagnosis of pure DCIS is made on NCB, but an associated invasive component is identified in the subsequent surgical excision specimen, raising the possibility of lymph node metastasis. If lymph node biopsy is not performed at the time of the definitive operative procedure of these cases, patients found to have an invasive cancer will require a second operative staging procedure. For example, in a previous study of 105 cases of pure DCIS diagnosed by NCB, Yen et al. found that 10% of these patients had a positive sentinel lymph node.

The invasive component may be underestimated on NCB with a diagnosis of pure DCIS in 15–20% of cases. In addition, Mendez et al. have identified microinvasive carcinomas in 8% of cases (6/77) diagnosed as pure DCIS on NCB. The number of DCIS upgraded to invasive carcinomas in surgical biopsies may be related to a variety of histopathological and radiological reasons. This upgrading of DCIS is more likely where there is high-grade DCIS, presence of periductal inflammation or lobular cancerisation, increasing number of calcifications, comedonecrosis with a cribriform/papillary pattern, and NCB from a palpable mass or large-sized lesions. DCIS underestimation rates also appeared to be in inverse proportion to the total volume of tissue examined (number of cores and needle gauge size). It is important when making a diagnosis of DCIS in NCB from a palpable mass to remember that most breast tumours exceeding 1.0 cm in size are at least partially invasive, especially when they are palpable. In our experience the underestimation of presence of invasion can be reduced by use of ultrasound guided biopsy targeting mass lesions if present and by use of multiple cores samples or larger volume vacuum assisted tissue sampling methods. This approach is supported by data indicating that the level of underestimation may be affected by the gauge of the needle and it has been suggested that this incidence can be reduced by using a thicker needle core. Lee et al. have reported that an 11-gauge vacuum-assisted biopsy was as reliable as open surgical biopsy for diagnosing DCIS without invasion. Previous studies have reported that the incidence of DCIS underestimates was 20–30% using a 14-gauge core, and using 4–10% with an 11-gauge core, and 4% using 8-gauge stereotactic vacuum-assisted biopsy. It is worth mentioning, however, that some authors could not identify any factor capable of predicting a higher likelihood of an invasive focus in surgical specimens following a diagnosis of DCIS by NCB, and recommended that patients with DCIS diagnosed by core biopsy should be offered sentinel lymph node biopsy.

It has also been reported that a smaller lesion size may be associated with increased risk for a negative or non-diagnostic core in patients with DCIS; the main reason for a false positive diagnosis of DCIS in NCB is atypical ductal hyperplasia (14–30% of cases), and in particular if the suspected lesion is restricted to one or two foci. On the other hand, most cases of DCIS in NCB which are suspicious or positive for microinvasion will demonstrate invasion in the subsequent excision. Thus, these cases should accordingly be carefully evaluated and we would recommend diligent examination of the core biopsy sample at multiple levels in such cases; if a definite diagnosis cannot be made, a surgical excision is recommended. In fact, the underestimation of malignancy in NCB and the role of atypical ductal hyperplasia is best reviewed in the article published Housami et al., who concluded, for instance, that a reliable diagnosis of atypical ductal hyperplasia may not be possible with NCB, because criteria for its diagnosis are based on surgical resection specimens.

After a diagnosis of DCIS is made on NCB, several other related variables should be addressed, such as extent of DCIS and its relative proportion to invasive component, nuclear grade, architecture, and the presence of necrosis and associated calcification. The combination of nuclear grade and presence of necrosis is currently the best predictor of biological behaviour of DCIS. Assessment of the presence and extent of DCIS component in a small invasive carcinoma is particularly important for breast conservation as patients whose tumours are predominantly in-situ, or which contain a significant in-situ component (>25%) and with evidence of DCIS outside of the invasive growth, are defined as having extensive intraductal carcinoma, a status that can adversely affect local control if the...
in-situ component is not adequately resected. Some of these patients with otherwise small carcinomas favourable for breast conservative therapy will not be candidates for breast conservation.21 In a previous study of DCIS in clinically palpable T3 and larger T2 carcinomas, El-Tamer et al40 reviewed whether NCB can identify patients with purely or predominantly in-situ carcinomas. They showed that NCB can reliably determine whether a palpable tumour mass is predominantly invasive or in-situ and accurately identify the probable extent of DCIS within an invasive mass. The authors also demonstrated a very high level of concordance between histological subtype, tumour grade, and the percentage of DCIS present in NCB and subsequent surgical biopsies.

Grading and histological typing of DCIS on core biopsy can be performed and is reasonably accurate. Current evidence suggests that concordance between DCIS grade on NCB and that in the definitive excision specimen is slightly similar to that of invasive tumours. In our institution, we found a concordance rate of 65% for DCIS grade compared to 67% for invasive tumour grade.40 Others have reported slightly better agreement for DCIS grade than for invasive grade.56

MICROINVASIVE CARCINOMA

Although minimally invasive “microinvasive” carcinoma (<0.1 cm) of the breast is a well-known and well-characterised entity in excision specimens, it is a rare finding in breast NCB and may require relatively extensive sampling to identify. Microinvasive carcinoma in NCB is commonly associated with DCIS and is often associated with larger invasive foci at excision. In a large series of NCB, Renshaw95 identified microinvasive carcinoma in 1% of cases (16/8500). The surgical specimens of these cases showed invasive carcinomas in 11 cases (2 with positive lymph nodes) and DCIS alone in 4 cases. Therefore, the term “microinvasive” carcinoma, like “DCIS”, should be used with care on NCB, and clinicians should be aware that there is a high probability of underdiagnosis of invasion.

LYMPHOVASCULAR INVASION

Lymphovascular invasion (VI) is another important prognostic factor in breast cancer. The presence of histologically confirmed VI is associated with an increased risk of metastatic disease, increased risk of local recurrence, and reduced survival.45 58 Some authors reported the inability of NCB to detect VI accurately.45 58 For example, Sharifi et al47 found that none of the 17 cancers with VI in the excision specimen showed VI on the NCB. However, in a study of 500 cases of invasive carcinoma diagnosed by NCB, we found that the concordance rate of detection of VI between NCB and surgical specimens was 69%.40 Our result was supported by the study of Pu et al,59 who assessed VI in the preoperative NCB to measure the pathological response after neoadjuvant chemotherapy; they were able to detect VI in NCB and found that its presence in NCB was associated with axillary lymph node metastases. They suggested that the presence of VI in NCB is an important feature that should be included in the standard report. In practical terms, when VI is identified in a needle core biopsy sample it should be reported as it may influence surgical management. There is, however, a significant failure rate of identification of at least 30% due to its microfocal nature.

TUMOUR NECROSIS

Assessment of necrosis in NCB may be useful in at least two settings: (1) intraluminal necrosis is one of the main features for diagnosis and grading of DCIS and is a pathological predictive factor for recurrence26 97 98; (2) assessment of necrosis can also help to predict and monitor pathological response to neoadjuvant chemotherapy.98

LYMPH NODE STAGE

It is well established that the lymph node stage is one of the most important prognostic factors for patients with breast cancer. Although there is little available data, in a previous study that included 166 patients presenting with primary operable breast cancer, we have shown that ultrasound-guided NCB and routine sectioning can identify lymph node disease in a relatively high percentage of cases (80% (27/34) of the sampled abnormal nodes).99 In a more recent study, NCB of the sentinel lymph node has been shown to be as accurate as excisional biopsy in detecting lymph node positivity by routine serial sectioning and H&E staining.100

BIOMARKER ASSESSMENT

In addition to histological information, NCB can be used to study the status of molecular markers with prognostic and predictive value that can contribute to an optimal selection of treatment strategy, especially for patients who may undergo preoperative therapy. The assessment of hormone receptors (HR) can be performed on NCB and correlates well with subsequent surgical excision specimens64 101 (table 1). The concordance rate was found to be higher for ER than PgR, which could be due to the fact that ER are more homogeneously distributed.65 102 103 It has been reported that HR scores correlated best in specimens that stain strongly and moderately positive; most discrepancies were between weakly positive and negative cases. Similar findings were reported in surgical specimens in which the major area of difficulty of reproducibility of HR staining was the group with low expression. In such cases, there is a risk of false negative results, mainly due to inadequate IHC assay sensitivity.104 When comparisons are made, NCB tends to up-score the receptor stain intensity compared with the subsequent surgical specimen, and in some cases, the cores show positive staining, while the surgical specimens are negative.51 104 105 Therefore, HR assays on NCB appears to be more reliable than assays on surgical specimens; better fixation on the core is the most likely explanation. Some authors, therefore, recommend that IHC for HR should be performed on core biopsy specimens, to ensure that patients with HR positive cancers receive appropriate hormonal therapy and are not overtreated with systemic chemotherapy.51 In our institution, we routinely assess ER in the core biopsy for invasive carcinomas. We repeat staining in the surgical specimen if the core biopsy shows weak staining or if the tumour shows morphological heterogeneity in the surgical resection and was ER negative on the core biopsy.106 HER2 status is recognised in breast carcinoma as a prognostic marker; overexpression is associated with worse prognosis. More importantly, testing is now required as a prerequisite to predict drug response and to select patients for trastuzumab treatment. Several reports have shown that the assessment of HER2 status on NCB in breast cancer is accurate and reliable.44 107 108 109 In a previous study of 325 patients with primary breast cancer, Taucher et al105 investigated the accuracy of HER2 status using IHC and fluorescence in-situ hybridisation (FISH) on NCB and compared the result with surgically obtained specimens. They found that the accuracy of IHC assessment of HER2 in NCB was 92% and increased to 96% with additional FISH analysis, which was applied to strongly positive cases. Therefore, they recommended performing FISH analysis for cases with strong IHC positivity on NCB in order to minimise the number of false-positive results.

In addition to HR and HER2 outlined above, a wide range of other molecular markers that have been identified in breast cancer can be assessed on NCB. Moreover, due to the relatively limited size of the specimens, NCB can help to determine the percentage of DCIS and invasive components in a tissue sample. This information can be valuable for determining the appropriate treatment strategy, especially for patients who may undergo preoperative therapy.
cancer, can be reliably assessed on NCB. For example, p53 and bcl2 have been evaluated on NCB; their assessment was proved to be accurate and reliable with a concordance rate of 85–100% when compared to surgical specimens. The accuracy of these markers was reported to be higher when they were scored as dichotomous variables (positive vs negative expression) rather than as continuous variables, possibly due to intratumoural cell heterogeneity. In addition, other markers can be reliably applied on NCB, such as E-cadherin to distinguish lobular from ductal carcinomas on NCB, myoepithelial markers (eg CK5/6, CK14, p63, CD10 or SMA), to differentiate between in-situ and invasive lesion. Interestingly, assessment of E-cadherin on NCB may also be relevant from the radiological perspective, as there are some indications that preoperative magnetic resonance imaging evaluation is accurate in determining true tumour extent in invasive lobular carcinoma, thereby impacting on the extent of therapeutic surgery.

OTHER ANCILLARY STUDIES
Estimation of microvessel density (MD) on NCB specimens in breast cancer may predict response to systemic therapy. Two previous studies have tried to assess MD on NCB and compare it with the subsequently excised tumours; however, both found that MD may not be reliably assessed on NCB and concluded that its usefulness, when measured as a continuous variable on NCB, is limited.

Gene expression data obtained from preoperative NCB of breast cancer patients can be used to build accurate predictive models that separate different molecular profiles. Several studies have shown the potential usefulness and reliability of NCB for high-throughput gene expression studies, such as semi-quantitative RT PCR, microarray technology and microRNAs (miRNAs) expression profiling. In addition, DNA analysis by flow cytometry was performed on NCB and the concordance with resected samples was high.

NEGATIVE SURGICAL EXCISION AFTER NCB
Another important point, which merits mention in this review, is the pathological approach when NCB removes the entire tumour, and the therapeutic surgical excision is devoid of residual tumour. Although no guidelines are available so far, we consider the best approach is to make sure that, in the surgical specimen, the core biopsy sample site is present, has been completely sampled and is completely removed. If these criteria are satisfied, then it is safe to state that the invasive component (or in-situ lesion) has been removed by the NCB sampling process. In this circumstance, the characteristics of the tumour should be assessed on the needle core sample—size, grade, type, VI, ER, PgR, and HER2. Tumour size should be correlated with the imaging size and it may be prudent to record the imaging size as the true size of the lesion.

CONCLUSION
NCB can reliably provide useful preoperative prognostic and predictive information in breast cancer patients which can play a major role in planning treatment strategies. Presence of invasive tumour, tumour type, histological grade, presence and extent of DCIS (including its features such as architecture type and grade) and tumour necrosis and lymph node status, if sampled, can be assessed relatively reliably on NCB. Assessment of prognostic and predictive biomarkers such as ER, PgR, HER2, MIB-1 and bcl2 and other ancillary tests can be performed on NCB and gives highly accurate results. Evaluation of vascular invasion, and minimal tumour invasion associated with DCIS may be less reliable. Primary tumour size and MD measurement may give unreliable values and their assessment on surgical specimens should be considered. However, it is important to emphasise the following: (i) meticulous operator technique is essential to provide representative samples; (ii) careful interpretation with prior knowledge of the limitations of the technique; and (iii) adherence to principles of triple assessment following biopsy can improve the diagnostic accuracy of NCB.

Take-home messages
- Needle core biopsy (NCB) is now considered the method of choice for tissue sampling as part of the triple assessment of breast disease.
- NCB can reliably provide useful preoperative prognostic and predictive information in patients with breast cancer, which can play a significant role in planning treatment strategies.
- Presence of invasive tumour, histological tumour type, presence of ductal carcinoma in-situ, hormone receptor status and HER2 overexpression can be assessed reliably on NCB.
- Meticulous operator technique is essential to provide representative samples.
- Careful interpretation with prior knowledge of the limitations of the technique and adherence to principles of triple assessment following biopsy can improve the diagnostic accuracy of NCB.

REFERENCES

AUTHORS' AFFILIATIONS

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