Aims: To establish the relation between the amount of breast core needle biopsy (CNB) material examined and agreement between preoperative and postoperative histopathology parameters in invasive breast cancer.

Methods: The CNB and surgical specimen histopathology reports of 113 patients with invasive breast carcinoma were reviewed and the total amount of CNB material examined for each case was determined. Agreement was calculated for tumour type, grade, mitoses, nuclear pleomorphism, and tubule formation. Associations between the amount of CNB material and histopathology agreement before and after surgery were explored using binary logistic regression.

Results: Tumour type and grade agreed in 65.4% and 61.6% of cases, respectively. The components used to calculate grade—nuclear pleomorphism (57.4%), mitoses (59.4%), and tubule formation (55.6%)—agreed slightly less frequently. The proportion of cases with preoperative and postoperative assessments that agreed did not depend on the number of cores collected or the total amount of material examined.

Conclusion: Neither tumour type and grade, nor the individual components used to calculate grade agreed consistently between the CNB and surgical specimen. The number of cores collected and the total amount of material reviewed by the pathologist does not influence the likelihood of agreement between preoperative and postoperative histopathology reports.

Breast cancer tumour sampling by core needle biopsy (CNB) is widely used in many centres and has proved to be up to 97% accurate at confirming the presence of malignant tissue within a suspicious lesion. It has been established previously that CNBs generally underestimate tumour grade and can also incorrectly type tumours (for review see Denley and colleagues). However, to our knowledge, there are no studies that have investigated whether the amount of material available to the pathologist from the CNB influences the agreement between the preoperative and postoperative histopathology of invasive disease.

Detailed preoperative histopathological information is increasingly required from pathologists and is potentially useful in determining the treatment of patients who may benefit from neoadjuvant chemotherapy, and also in the development of preoperative prognostic models. The ability to determine accurately tumour type and grade and therefore the individual components of the grade—tubule formation, mitoses, and nuclear pleomorphism—is crucial in the development of any preoperative prognostic tool. It seems probable that there is a correlation between the amount of clinical material obtained from the CNB and its accuracy. Therefore, we explored the relation between the amount of clinical material available for examination by the pathologist and how faithfully preoperative histopathology reflects that of the surgical specimen.

“Detailed preoperative histopathological information is increasingly required from pathologists”

Patients and Methods

The pathology records at our centre from 1999 to 2002 were reviewed to identify those patients who had undergone CNB for a suspicious breast lesion. Patients were included in our study if both their CNB and surgical specimen reports were unequivocal in describing the presence of invasive breast carcinoma. Patients who had previously received, or were receiving, chemotherapy were excluded, and 113 patients who had undergone CNB (all Tru-cut 1 biopsy needle) and subsequent primary surgical excision for breast carcinoma were identified. All patients had been managed in accordance with the UK National Health Service (NHS) breast screening programme guidelines, which require surgical excision within two weeks of diagnosis. The original haematoxylin and eosin stained CNB slides were reviewed and retrieved by a consultant histopathologist, and we collected the following information from the histopathology reports: anonymised patient identification number, type according to the UK NHS breast screening programme guidelines, grade, nuclear pleomorphism, mitoses, and tubule formation according Elston and Ellis’s modification of the Bloom and Richardson system.

Core length was measured manually using a ruler, but to ensure the accuracy of this method at detecting and measuring all of the biopsy material on the slide we took a random sample of 10% of the biopsies and measured these using the Nikon Eclipse E-1000 microscope/Lucia image analysis system (Nikon, Kingston upon Thames, UK). No significant difference between the two methods of measurement was found (p = 0.421, paired t test). A “core index” for each patient was calculated, which was derived by multiplying the number of cores taken by the number of levels viewed by the histopathologist by the total length of the cores. For example, for three levels of three cores measuring 23 mm in total the core index would be 207. Repeated levels were occasionally requested for cores where there was difficulty in confirming the diagnosis. Potentially, this could confound our analysis, so that cases where only the

Abbreviations: CNB, core needle biopsy; NHS, National Health Service.
standard three levels were available (n = 110) were examined as a separate dataset. Binary logistic regression (using MinitabTM Statistical Software 13.1; Minitab Inc, State College, Pennsylvania, USA) was used to assess our hypothesis that the proportion of preoperative and postoperative assessments that agree may be a function of the number of CNBs collected or the total amount of clinical material obtained. Two models, one containing the number of cores and one with the core index as an independent variable, were investigated for each histopathology variable. Independent variables were assessed at the 10% level of significance.

RESULTS
Agreement between the CNB and the surgical biopsy specimen for grade, mitoses, tubules, pleomorphism, and type in 113 patients with breast cancer was assessed. Tumour type agreed in 65.41% of cases, which was similar to grade (61.65%). The components used to calculate grade—nuclear pleomorphism (57.40%), mitoses (59.40%), and tubule formation (55.64%)—agreed less frequently. Grade and mitoses were more likely to be underscored preoperatively than overscored; grade was underscored preoperatively in 24.06% of cases compared with 14.28% overscored, and mitoses were underscored in 33.08% of cases compared with 7.52% overscored. Tubule formation was more likely to be overscored preoperatively (12.78%) than postoperatively (31.58%). Pleomorphism was overscored and underscored equally frequently (21.05% ± 21.81%).

Table 1 shows the assessment of agreement between preoperative and postoperative assessments for tumour type, grade, mitoses, nuclear pleomorphism, and tubules. The number of cores did not influence the proportion of cases with agreement between CNB and postoperative histopathology (p = 0.42, binary logistic regression). The more sophisticated measure of the amount of clinical material viewed by the pathologist, the core index, was used to investigate further the influence of clinical material on agreement. This was calculated from the number of cores, total length, and number of levels viewed, and was therefore effectively an index of the information available to the pathologist. Again, using binary logistic regression, there was no significant association between the amount of biopsy material viewed and the proportion of preoperative and postoperative histopathology agreements. We were concerned that our results could be compromised by the cases where repeated levels were requested. Therefore, we examined the 110 “standard” cases where only levels 1–3 were viewed as a separate dataset. Again, no significant difference was found in either the mean number of cores or the mean core indices for agreement between the parameters (all p values > 0.1, binary logistic regression).

The means, medians, ranges, and distributions of core indexes for grade and tumour type were, in fact, almost identical between the two groups shown in fig 1, which describes the full dataset of 113 patients. The distributions of core indexes for mitoses, pleomorphism, and tubule formation are not shown, but were broadly similar, as would be expected from the results for tumour grade.

DISCUSSION
Accurate preoperative assessment of breast cancer is becoming increasingly important. Surgeons and pathologists are looking beyond the ability to diagnose invasive disease accurately to tools that will allow the earlier selection of patients for adjuvant chemotherapy or for more conservative surgical procedures.

We have assessed tumour grade, tumour type, and the components used to calculate grade. We used the grade components for two reasons: (1) if grade did not frequently agree with postoperative histology we wanted to establish whether one particular component contributed disproportionately to the disagreement; and (2) perhaps more importantly, we were interested to see whether we could identify one component that was consistently accurate, and which would therefore make a more suitable candidate for preoperative prognosis.

Broadly, we found that all of the parameters we assessed agreed between preoperative and postoperative histopathology in 55–65% of cases. This is consistent with two similar studies. However Harris et al reported significantly higher rates of agreement for tubule formation and pleomorphism than was seen in our study. Where there was disagreement, it was more likely to be underscored preoperatively than overscored, which is a well recognised phenomenon. As other authorities have proposed, it is probable that the grading of the surgical specimen, which (by necessity) relies on finding the worst areas to score the grade components, will generally grade higher than a CNB, which samples a very small fraction of typically heterogeneous tumours.

<table>
<thead>
<tr>
<th>Number of cores</th>
<th>Core index</th>
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<tr>
<td>N</td>
<td>Median</td>
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<tr>
<td>Type</td>
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<td>Grade</td>
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<td>Mitoses</td>
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<td>Mitoses</td>
<td>Disagree</td>
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<td>Tubule formation</td>
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<td>Tubule formation</td>
<td>Disagree</td>
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<td>Pleomorphism</td>
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<td>Pleomorphism</td>
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In total, 113 patients were examined to find out whether the number of cores collected during the diagnosis of a suspicious lesion correlated with the likelihood of agreement between preoperative and postoperative histology. This was assessed by two methods: (1) a count of the number of cores and (2) an index of the amount of material examined by the pathologist (as described in Patients and methods). Mitoses, tubule formation, and nuclear pleomorphism were components of tumour grade. The p value was determined by binary logistic regression.
In cases of invasive breast carcinoma, the number of cores and the amount of material viewed by the pathologist does not influence agreement between the histopathology reported from the preoperative core biopsy and the definitive excision specimen.

**Take home message**

- In cases of invasive breast carcinoma, the number of cores and the amount of material viewed by the pathologist does not influence agreement between the histopathology reported from the preoperative core biopsy and the definitive excision specimen.

We have asked a fundamentally different question to previous work examining the relation between the number of cores and diagnosis. Typically, these studies have sought to determine the relation between CNB number and diagnostic accuracy. We were interested to find out whether, in cases where there is a definite diagnosis, the amount of clinical material available influences agreement between CNB and the surgical specimen.

"This surprising and somewhat paradoxical result defies simple explanation"

We could find no significant association between either the number of cores collected or the total amount of material viewed by the pathologist and the likelihood of agreement. This surprising and somewhat paradoxical result defies simple explanation. It would be expected that the more material sampled from a tumour the better the representation and therefore agreement. One of many possible explanations is that doubling or even tripling of the number or length of a typical CNB only represents single per cent sampling increases from the total volume of even a 1 cm³ tumour.

However, these data do suggest that provided there is sufficient CNB to confirm the presence of invasive disease, there is little to be gained from further biopsies in terms of both accurate staging of grade and tumour type and also improving the quality of information available for putative prognostic tools. Moreover, we show that confidence in the CNB data to provide prognostic information does not increase with larger or more biopsies.

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