Comparison of fine needle aspiration cytology and needle core biopsy in the diagnosis of radiologically detected abdominal lesions

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Aims: To compare the sensitivity and specificity of percutaneous fine needle aspiration (FNA) cytology and needle core biopsy (NCB) in the diagnosis of suspected intra-abdominal tumours.

Methods: One hundred and forty one consecutive patients who underwent radiologically guided combined FNA/NCB of abdominal lesions over a four year period were reviewed. The diagnostic accuracy of both techniques and the value of rapid staining and assessment of cytological preparations were assessed.

Results: FNA cytology and NCB identified 111 of 129 (86%) and 104 of 129 (80.6%) malignant lesions, respectively; in combination, the sensitivity increased to 90.7%. The diagnostic specificity was 100% for both methods, although one case of phaeochromocytoma was misinterpreted as undifferentiated carcinoma on biopsy. More accurate tumour subtyping was possible in two cases with FNA and four cases on NCB. The series included 12 benign lesions, of which 11 and nine were accurately identified on FNA and NCB, respectively. Two specific benign diagnoses (Budd-Chiari syndrome and hepatic infarct) were made only on biopsy. The use of rapid assessment cytology preparations ensured that appropriate samples were submitted for microbiology in three liver abscesses, and provided an accurate cytological diagnosis at the time of the procedure in 103 of 141 (73%) cases. None of the patients suffered biopsy related complications.

Conclusions: FNA cytology is more sensitive and accurate than NCB in the diagnosis of abdominal lesions, and also offers more rapid diagnosis. However, the combination of these sampling techniques increases diagnostic sensitivity and occasionally provides more accurate classification of tumours and benign lesions. The techniques should be considered complementary in the investigation of abdominal lesions.

The management of patients with suspected neoplastic disease involving abdominal sites is dependent on obtaining an accurate tissue diagnosis, usually via percutaneous sampling. Patients with abdominal lesions may present with clinically evident tumour masses, but the increasing use and sensitivity of radiological techniques has also led to the identification of relatively small lesions, which require the use of image guidance for reliable targeting. At present, there are two widely used and accepted methods for obtaining diagnostic material, namely fine needle aspiration (FNA) cytology and needle core biopsy (NCB). FNA specimens are usually acquired using 20–25 gauge needles and generally provide a sample for cytological examination, whereas NCB specimens are obtained using larger 14–18 gauge needles and primarily provide a tissue core for histological assessment. In theory, each sampling method offers different advantages and limitations. Although both techniques are very safe, FNA is often preferred in sampling deeply placed lesions, sites adjacent to major vessels, or in situations in which needles are to be passed through the bowel wall. Cytological samples can be rapidly stained and examined, thereby providing immediate assessment of adequacy, and in many cases a provisional diagnosis can be made while the patient remains in the radiology department. Furthermore, involvement by pathologists on site optimises clinical correlation and ensures that specimens are optimally handled and that appropriate samples are taken as required for ancillary investigations, such as microbiology or molecular studies. The advantages of NCB include the greater familiarity of histological preparations among some pathologists, the preservation of tissue architecture, which may be important in the assessment and subtyping of some tumours, and the relative ease with which histochemical and immunohistochemical techniques can be applied to paraffin wax embedded biopsy material.

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Therefore, although it might appear that cytological and histological examination would be complementary in the assessment of abdominal lesions, there are conflicting data in the literature regarding the accuracy and usefulness of these techniques. In particular, there are wide variations in the reported diagnostic sensitivities of FNA cytology and NCB, to the extent that some authors have suggested that core biopsy alone should be used, or that FNA is the preferred technique with biopsy limited to cytologically indeterminate cases. These discrepancies may be partly explained by variations in the types of lesion subject to biopsy, and by the approach to cytological examination (such as the use of rapid staining techniques). Relatively few reports have evaluated FNA cytology and core biopsy obtained in combination in the investigation of patients with abdominal lesions. Therefore, we have compared the sensitivity and specificity of FNA cytology and...

Abbreviations: FNA, fine needle aspiration; NCB, needle core biopsy; NOS, not otherwise specified
PATIENTS AND METHODS

The histopathology and cytopathology databases of the pathology department, Glasgow Royal Infirmary were searched for all patients undergoing combined, image guided percutaneous core biopsy and FNA cytology sampling of abdominal mass lesions during the four year period August 1995 to July 1999. All patients had one or more radiologically detected lesions and the initial clinical suspicion in each case was of neoplastic disease. In total, 141 cases were identified, comprising approximately 25% of all abdominal lesions subject to biopsy in the study period (in most cases FNA or NCB alone was performed). There were 76 men and 65 women with an age range 17–88 years (mean, 65.7). Eighteen patients had a previous history of biopsy confirmed malignancy as follows: carcinomas of the colon (n = 9), stomach (n = 4), breast (n = 2), and pancreas (n = 2), and ocular malignant melanoma (n = 1). Coagulation screen was checked before the procedure. In general, FNA samples were performed following local anaesthesia using a standard 21 gauge Chiba needle attached to a 20 ml syringe. After localisation, the needle was passed gently through the lesion four to six times with aspiration. The needle was withdrawn and passed to the cytopathologist. Direct smears were prepared from each sample and one or more slides were stained using the Diff-Quik method in the scanning room. The smears were examined by the cytopathologist and the radiologist was informed of specimen adequacy. Repeat FNA samples (up to three in total) were taken as required if limited material was obtained. A provisional diagnosis was made whenever possible and in some cases this was included with the radiology report returned to the ward with the patient. After preparing smears, the needles were rinsed in normal saline. Cytospin or cell block preparations were made in a few cases for subsequent ancillary investigations, such as histochemistry or immunocytochemistry, most commonly to confirm the epithelial nature of poorly differentiated malignancies or to demonstrate lymphoid or mesenchymal antigens. The needle rinses were also submitted for microbiological study in those cases suspected to be of inflammatory or infective nature on rapid cytological assessment. After the FNA procedure, core biopsy samples were taken using an 18 gauge needle loaded into an automatic biopsy system (Biopry System, Radiplast, Sweden). The adequacy of the specimens was judged visually and up to three separate core samples were taken as required. The core biopsies were fixed in 10% neutral buffered formalin, processed routinely, and stained with haematoxylin and eosin. The specimens were examined and reported as part of the normal diagnostic workload by cytopathology and histopathology staff.

For the purposes of our study, the histological diagnoses from NCB specimens were considered the “gold standard”. In those patients with negative or unsatisfactory core biopsy samples the definitive clinicopathological diagnosis was based on subsequent biopsy or surgical resection specimens, or on clinical follow up data obtained by case record review.

RESULTS

Table 1 outlines the final clinicopathological diagnoses for the 141 patients. One hundred and twenty nine patients had malignant disease (two adrenal phaeochromocytomas are included in the potentially malignant group). The most common diagnosis was metastatic malignancy in the liver (87 cases), comprising 82 metastatic carcinomas, three lymphomas and one case each of melanoma and sarcoma. There were nine cases of primary hepatocellular carcinoma, all of which were diagnosed on FNA; in one case, the NCB showed only cirrhotic liver tissue (fig 1). All pancreatic and renal tumours were primary carcinomas at these sites. The retroperitoneal tumours comprised two leiomyosarcomas and one case of Hodgkin’s disease. The miscellaneous abdominal/pelvic tumours comprised seven patients with peritoneal metastatic adenocarcinoma, one case of borderline mucinous tumour of the ovary, and one patient with a malignant gastrointestinal stromal tumour of uncertain origin.

In three patients both the FNA samples and the core biopsies were inadequate for assessment (acellular or lacking parenchymal elements); subsequent follow up showed metastatic adenocarcinoma in the liver in two cases and adrenal phaeochromocytoma in one. Three further patients had unsatisfactory NCB specimens but positive cytology, whereas...
two patients had inadequate FNA but diagnostic core biopsy. Altogether, six NCB and five FNA specimens were inadequate. In the remaining false negative cases, parenchymal tissue was obtained on sampling but later shown to be unrepresentative of the primary lesion. Overall, FNA cytology accurately identified 111 of 129 malignancies (86%), whereas 104 cases (80.6%) were positive on core biopsy. Combining the techniques increased the sensitivity to 90.7% for patients with malignant disease.

There were relatively minor discrepancies between the cytological and histological diagnoses in seven of the malignant cases, all involving liver specimens. In two cases, the FNA samples were reported as consistent with metastatic adenocarcinoma, whereas the corresponding core biopsies showed a large cell carcinoma lacking specific differentiation. Conversely, two large cell carcinomas that were not otherwise specified (NOS) on cytology showed evidence of glandular differentiation on core biopsy. One case was reported as malignant, NOS, on FNA because there was insufficient material for ancillary studies; the corresponding biopsy showed an undifferentiated carcinoma and this was supported by cytokeratin immunoreactivity in the tumour cells. In one case, both the cytology and biopsy showed metastatic adenocarcinoma but the histological pattern was suspicious of prostatic origin and this was confirmed using immunostaining for prostatic acid phosphatase and prostate specific antigen. Finally, one case was reported as favouring primary hepatocellular carcinoma on FNA, whereas metastatic adenocarcinoma was considered more likely on core biopsy; the patient died without resolution of the diagnosis.

There was an additional tumour in which the histological diagnoses on core biopsy and on the subsequent resection specimen differed. This was an adrenal mass in which the core biopsy was interpreted as undifferentiated carcinoma but the surgical specimen showed a phaeochromocytoma. There was no evidence of metastatic disease at the time of surgery. The FNA sample was unsatisfactory in this case and the diagnosis was made on tumour resection. There were 11 inflammatory/non-neoplastic lesions comprising three liver abscesses, one liver infarct, one case of Budd-Chiari syndrome, four non-specific reactive liver patterns, and two cases of pancreatitis. The FNA samples were adequate in all cases and were reported as consistent with inflammatory/reactive processes. Material from the FNA samples was submitted to microbiology in six cases and Streptococcus milleri was cultured from two of three hepatic abscesses.

Core biopsy samples from two of the inflammatory lesions (one pancreatitis, one non-specific liver reaction) were inadequate for diagnosis. The diagnoses of Budd-Chiari syndrome and hepatic infarct were based on the core biopsy findings because the cytology specimens showed non-specific liver tissue.

Overall, therefore, FNA cytology and core biopsy were diagnostic in 122 of 141 (86.5%) and 113 of 141 (80.1%) cases, respectively. A combination of cytology and biopsy improved the diagnostic yield to 90.8% of cases (128 of 141).

A provisional diagnosis was made at the time of the procedure in 103 FNA (73%) specimens (92 malignant cases and 11 reactive/inflammatory lesions). There was no difference between the provisional and final cytology diagnoses in these cases. The definitive cytology report was issued within 24 hours of the procedure in 74% of cases and the mean reporting time was 1.4 days (range, 0–8). The NCB reports took 3.9 days on average to complete (range, 1–12).

None of the patients in the series had serious complications after the biopsy procedures.

**DISCUSSION**

Needle biopsy of abdominal mass lesions may be used to establish a malignant diagnosis in patients with clinically or radiologically suspected neoplasia or for staging in patients with known tumours at other sites. The decision to use FNA and/or NCB as sampling techniques depends on many factors including the size and site of the lesion, the suspected likely diagnosis, and the risk of complications. Because most biopsies are performed using image guidance, the experience of individual radiologists is an

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**Figure 1** Hepatocellular carcinoma. (A) Fine needle aspiration sample includes thickened trabeculae of hepatocytes with peripheral endothelial cells (arrows) and scattered atypical bare nuclei (above). Diff-Quik method, original magnification, ×300. (B) The corresponding core biopsy shows cirrhotic liver but no evidence of malignancy. Haematoxylin and eosin, original magnification, ×120.
important factor, and the preferred technique may be influenced by the availability of cytopathologists for on site specimen assessment. The sensitivity and specificity of FNA and NCB should also be considered in choosing the optimal technique.

FNA proved more sensitive than NCB in the diagnosis of the 129 malignant lesions in our series (86% v 80.6%, respectively). Similar findings have been recorded in most previous studies in which cytological and histological sampling of abdominal masses have been directly compared (table 10–12 14–19). The sensitivity of FNA was 2–24% greater than that of NCB in these studies, although some reports excluded unsatisfactory specimens in their analysis.11 As with previous reports, we found that the combination of cytology and core biopsy increased the sensitivity of the biopsy procedure. However Nyman and colleagues10 reported only 61.8% sensitivity for FNA compared with 90.1% for NCB in the investigation of 55 patients with malignant liver lesions. The same group also reported broadly similar findings in the diagnosis of paediatric abdominal tumours.9 The authors concluded that NCB should be the preferred sampling technique in patients with abdominal masses and that the combination of FNA and NCB had no additional value. Moulton and Moore11 also found biopsy to be more sensitive than FNA in the assessment of lesions at various anatomical sites (86% v 75%, respectively in abdominal malignancies). However, the false negative rates for FNA in these studies were greater than those in most other combined FNA/NCB series, or in recent reports using FNA cytology alone, in which sensitivities over 90% have been recorded.20–23 It is also pertinent that neither group of authors used immediate assessment of cytology samples which we, and many others,1 7 feel maximise diagnostic yield and accuracy.

In our study, FNA cytology was 100% specific for a malignant diagnosis. Although rare false positive diagnoses have been described with FNA,1 1 2 5 2 6 2 8 most studies have confirmed that cytological examination is highly reliable for a malignant diagnosis in abdominal lesions. We also considered NCB to be 100% specific in the current series, although it could be argued that there was one false positive histological diagnosis. This case involved an adrenal mass that was interpreted as undifferentiated carcinoma on core biopsy, but subsequent tumour resection revealed a phaeochromocytoma without evidence of metastatic spread. Although we included the two phaeochromocytomas in our series within the malignant category, most such tumours are clinically benign and the NCB assessment in the above case could therefore be regarded as misleading and inaccurate.

"FNA cytology was 100% specific for a malignant diagnosis"
Fine needle aspiration cytology is more sensitive and accurate than needle core biopsy (NCB) in the diagnosis of abdominal lesions, and also offers more rapid diagnosis. However, NCB can provide specific tumour subtyping in a small number of cases. Therefore, these two techniques should be considered complementary in the investigation of abdominal lesions.

In conclusion, a direct comparison of FNA and NCB in 141 patients undergoing image guided sampling of abdominal lesions showed that FNA cytology was more sensitive and accurate than biopsy, and that rapid diagnosis was also usually possible with this technique. However, NCB offered the advantage of specific tumour subtyping in a small number of cases. We feel that FNA and NCB should be considered complementary diagnostic techniques and used in combination depending on clinical conditions.

**References**


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