Role of ultrasound guided fine needle aspiration biopsy in the diagnosis of hepatocellular carcinoma

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Abstract
In 170 cases of hepatocellular carcinoma, ultrasound showed a high sensitivity in identifying focal liver lesions. Fine needle aspiration biopsy guided by ultrasound yielded a pathological diagnosis in the majority of cases. The advantages of this technique, its high diagnostic yield and low cost, render the technique feasible for all units. Complementary use of fine needle aspiration biopsy under ultrasound guidance and laparoscopy assures the highest rate of diagnostic accuracy in hepatocellular carcinoma. We confirm the poor sensitivity of α-fetoprotein levels in distinguishing hepatocellular carcinoma from other liver neoplasms.

The incidence of hepatocellular carcinoma in some areas of Asia and Africa reaches 150 cases/100,000 inhabitants per year; in these areas hepatocellular carcinoma is the most common neoplasm. In Europe, the United States, and Australia the reported incidence is less than 5 cases/100,000. It is probably underestimated, however.

Hepatocellular carcinoma is more frequent in men and in the middle aged and elderly. But in the areas where hepatitis B is endemic, the prevalence of this tumour is higher and the average patient age lower. Known risk factors are hepatitis B virus, alcohol, and liver cirrhosis. α-Fetoprotein concentrations show poor sensitivity for early diagnosis of hepatocellular carcinoma. Detection is facilitated by ultrasonography, whereas pathological diagnosis is possible with guided fine needle biopsy. The diagnostic accuracy of cytology in hepatocellular carcinoma has now been confirmed by several studies.

Early diagnosis is important in order to modify the extremely poor prognosis of this tumour by surgical intervention. In this paper we report data from 170 consecutive patients with hepatocellular carcinoma seen in our department. The aim is to point out the diagnostic role of ultrasound guided fine needle aspiration biopsy in the diagnosis of hepatocellular carcinoma.

Methods
We report on 170 consecutive patients with hepatocellular carcinoma seen from June 1981 to February 1989. Hepatitis B virus markers (Enzymo Assay Laboratories and Enzymo Immunoassay Behring Institute) assays were performed, as well as routine liver function tests. In all patients abdominal ultrasonography was performed with real-time equipment (Hitachi EUB 22,26 and Aloka SSD 650 with 3-3.5 MHz linear or convex probes). In some cases computed tomography, selective liver angiography, and laparoscopy were carried out. Ultrasound guided fine needle aspiration biopsy (Chiba needle, 22 gauge) was performed according to the technique we previously described. Aspiration was performed with an aspirating cannula with short forward and backward movements visualising the bright echo of the needle tip in the target.

All patients had platelet counts of over 70 x 10^4/l and prothrombin times (% of the control) higher than 50% (international normalised ratio 1:3). A cytology sample, stained by a rapid May-Grunwald-Giemsa method (Merck Hemacolor), was examined immediately and a subsequent biopsy was performed only when the first specimen was inadequate. The other samples, stained with Papanicolaou and May-Grunwald-Giemsa, were examined by an experienced cytologist for definitive diagnosis.

In three cases of well differentiated hepatocellular carcinoma the cytological suspicion was confirmed by a cutting fine needle biopsy (Surecut). Verification of the cytological diagnosis was obtained by histological examination of samples obtained during laparoscopy (11 cases), surgery (eight cases), angiography (15 cases); by α-fetoprotein concentrations (>500 ng/ml) (35 cases); and by correlation with clinical, laboratory, and imaging data (63 cases).

The cytological diagnoses were categorised as true positives, true negatives, false positives, false negatives, and non-diagnostic (inadequate specimens/insufficient material). In this way we have evaluated the sensitivity, specificity, accuracy, and positive and negative predictive value of fine needle aspiration biopsy in hepatocellular carcinoma and calculated the significance of the results using the χ² test.

Results
The study group comprised 131 men and 39 women (male to female ratio 3:1). The mean age was 65.4 years (64.5 years for men and 68.8 years for women), range 40–83 years.

Liver cirrhosis was ascertained in 149 of 170 (87.6%). In 66 cases diagnosis was made by histology or laparoscopy, or both, and in the other cases by clinical, laboratory, and ultrasound data. Serum hepatitis B virus markers were measured in 160 of 170 cases and 22 patients were HBsAg positive (13%). Fetal protein was measured in 165 of 170 patients: normal concentrations were found in 63 cases (38%), 21–500 ng/ml in 45 cases (32%), and >500 ng/ml in 48 cases (29%). α-Fetoprotein concentrations did not vary significantly with
tumour size, cytology, or hepatitis B virus markers.

Ultrasonography was carried out in all patients: the liver lesion was detected in 167 of 170 (98%); the nature of the lesion was misinterpreted in three cases. The tumour appeared as a single nodule in 77 cases (the diameter was less than 5 cm in 34 cases (20-3%) and less than 3 cm in 16 cases (9.5%)); as multiple nodules in 51 cases, and was diffuse in 39 cases; 46% of the lesions with a diameter of less than 5 cm appeared hypoechoic.

Computed tomography, performed in 40 patients, detected lesions in 86% of the cases; angiography, carried out in 28 patients, showed 100% accuracy.

Altogether, 138 patients underwent ultrasound guided fine needle aspiration biopsy. The diagnosis of hepatocellular carcinoma was obtained in 132 cases (sensitivity 95.6%; specificity 100%); 68 cases showed a well differentiated cell tumour, in 50 the tumour was poorly differentiated, and 14 showed a large pleomorphic cell type. We report only one case of bleeding after biopsy, which required blood transfusion (500 ml).

In the other 38 patients (six fine needle biopsy false negatives and 32 remaining patients without fine needle biopsy) a definitive diagnosis was reached with ultrasound guided 2-1 mm trucut biopsy (one case); with laparoscopy and associated biopsy (13 cases); on the basis of clinical and laboratory data and ultrasonography (20 cases); and at necropsy (four cases).

In this series the 44 laparoscopies were performed to establish the initial diagnosis, to correct false negative fine needle biopsy, to determine presurgical staging, and to check the true positives of fine needle biopsy during its introductory phase into our department. The sensitivity of laparoscopy was 70-5%. The sensitivity was better when this technique was used to check the true positives of fine needle biopsy or to correct false negatives where laparoscopy was not performed in the cases with deeply located lesions.

Discussion

Laboratory diagnosis of hepatocellular carcinoma is principally based on $\alpha$-fetoprotein titres, even though many workers point out the poor sensitivity of this marker for early diagnosis.11-14 We found normal values in 38.2% of the cases and highly diagnostic values in a quarter of the patients. We did not find a significant correlation between $\alpha$-fetoprotein values and hepatocellular carcinoma cytology or tumour dimensions.

In the present series the percentage of the single hepatocellular carcinoma lesions with a diameter of less than 3 cm is small, 9.5%. In another Italian study13 the percentage was 10-5%.

To detect hepatocellular carcinoma we always performed ultrasonography, whose sensitivity is considered to be very good.15,16 For lesions of less than 2 cm it gave a diagnostic accuracy higher than computed tomography and similar to angiography.17 Recent reports (on a small series of hepatocellular carcinoma) attribute a higher diagnostic accuracy to computed tomography and magnetic resonance imaging than to ultrasonography. Ultrasonography must nevertheless be considered the primary technique for safety, economy, and good definition of the vascular and biliary trees.18,19 In addition, it can optimally guide the biopsy needle.

In selected cases ultrasonography with pulsed Doppler is useful to avoid haemorrhagic complications of fine needle aspiration biopsy guided by ultrasonography either by showing the vascular nature of the target lesion, or by showing a highly vascular area around the mass.20

Pathological diagnosis was reached in most of our patients with ultrasound guided fine needle biopsy, which showed a high diagnostic accuracy with a sensitivity of 95-6% in 138 patients. Bret et al21 used the procedure in 159 cases of hepatocellular carcinoma, obtaining a diagnostic sensitivity of 84% (92% for sensitivity in predicting the malignancy).

The cytological diagnosis of hepatocellular carcinoma can be difficult in well differentiated cell types. In fact, four of our six false negative cases were well differentiated cell type hepatocellular carcinoma (the other two cases were technically difficult to approach: small and deep target lesions). Recent reports cite a diagnostic accuracy of cytology (sampling under laparoscopic guide) ranging from 93% to 100%.22 Bret et al21 point out the usefulness of comparing hepatocytes sampled from tumour and from normal areas of the liver. Pedio et al20 have shown the importance of naked nuclei, which show the same alterations as intact tumour hepatocytes. The phenomenon seems to be characteristic of hepatocellular carcinoma and is absent in liver cirrhosis.

To improve diagnostic accuracy Limberg et al22 suggest the use of cutting fine needle biopsy. Their results, however, are similar to ours and to those of Bret et al.21 We have used cutting fine needle biopsy in only three cases of well differentiated hepatocellular carcinoma to correct an equivocal cytological ‘negative.’ Such a manoeuvre, however, increases the cost of the investigation and may increase the risk to the patient. In our series we had one case of bleeding. In the series of Bret et al21 there were four cases of bleeding, one of which was lethal.

In a recent series24 reporting the complications of abdominal fine needle biopsy, the mortality was 0-018% (the two deaths were due to haemoperitoneum in cases of hepatocellular carcinoma with cirrhosis). We agree with Hall-Craggs and Lees25 that the use of the cutting fine needle has to be reserved for those cases not diagnosed by cytology. We emphasise that the rapid staining of the material aspirated during ultrasound guided biopsy can reduce the number of biopsy specimens required from each patient without impairing the diagnostic accuracy26 and can thereby reduce the complication rate associated with the biopsy.

We point out that ultrasound guided fine needle biopsy should replace blind percutaneous biopsy22 for typing of hepatocellular carcinoma and other focal liver lesions. The introduction of fine needle biopsy has modified the indications for laparoscopy.22
For the initial diagnosis we used laparoscopy in selected cases: suspected hepatocellular carcinoma with negative or inconclusive ultrasound, lesions with ultrasound features resembling subcapsular haemangioma, or cases where there were difficulties in performing fine needle biopsy. Laparoscopy is advisable to correct false negatives and for presurgical staging. Moreover, it is useful for the study of equivocal lesions in a cirrhotic liver with adenomatous hyperplasia.8,9

We think that ultrasonography is a good method for identifying hepatocellular carcinoma, while ultrasound guided fine needle aspiration biopsy seems to be the first choice of the invasive technique. The indications for laparoscopy are thereby reduced, but it retains an essential role in selected cases. The integration of these two techniques gives excellent diagnostic accuracy.17

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