Advances in neoplastic disease of the liver and biliary tract

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The last 25 years have seen an explosion of interest in hepatic and biliary neoplasia reflecting not only the worldwide importance of these tumours, but also the fact that recent advances in serological and radiological diagnosis and the new techniques of molecular biology have all had a major impact. It seems hard to believe that in 1965, none of the hepatitis viruses had been identified and although 'Australian antigen' had just been described its significance was yet to be realised. Computerised tomography, ultrasonography and ERCP were all unheard of. The idea of useful tumour markers was still the stuff of oncologists’ dreams, alphafetoprotein having only been recognised in animal studies in the year the first liver transplant programme was started (1964). This review aims to summarise progress over the period in question and highlight some of the contributions and current areas of interest of the Institute over this period.

Historical overview

In 1965 there were occasional perceptive suggestions that hepatocellular carcinoma might be aetologically related to chronic infection with one of the hepatitis viruses. With the discovery of the association between 'Australia antigen' and hepatitis in 1967, however, the door was opened for a flood of papers soon to show an undoubted association of this tumour with hepatitis B virus infection.1 The crowning epidemiological study was that of Beasley et al who, in an ongoing follow up study of over 22 000 male Chinese, clearly showed that those who were HBsAg seropositive were at a relative risk of over 100 for the development of hepatocellular carcinoma.2 At about the same time there were several reports that integrated HBV–DNA could be detected in the genome of hepatoma cell lines and in some samples of human liver tumour tissue.3 Another seminal observation was that certain animals, particularly woodchucks4 and Pekin ducks carried viruses (subsequently classified as 'hepadna' viruses) which are similar in genome size, morphology, and genome organisation to human hepatitis B virus. The woodchuck frequently develops chronic hepatitis and hepatocellular carcinoma which contains integrated WHV–DNA sequences and is, therefore, an excellent model for the investigation of hepatitis B virus related carcinogenesis.

It is probably fair to say that molecular biological investigations made rather less progress than expected over the next decade, in explaining the mechanism of hepatitis B virus carcinogenesis. Some workers made the startling observation that even among alcoholic cirrhotic patients developing hepatocellular carcinoma and subjects with no serum markers of hepatitis B virus infection, integrated HBV–DNA could be detected in tumour tissue.4 These observations remain controversial, and others have been unable to confirm them.5 There is currently no evidence that the viral DNA contains a transforming gene — that is, an 'oncogene' — and analysis of integrated viral segments has failed to reveal any consistent site of integration or common deletions or duplications.

The excitement over hepatitis B virus involvement overshadowed other possible aetiological factors particularly cirrhosis, the frequency of which amongst men with hepatocellular carcinoma had been commented upon for at least 50 years.6 In a large prospective follow up of 613 patients with different types of cirrhosis initiated in 1978 by Walter Melia and continued by Shaams Zaman and Elizabeth Metivier, multivariate analysis showed that the major independent risk factors for the development of hepatocellular carcinoma were increasing age, 'non-UK' nationality and male sex.7 Interestingly, serum hepatitis B virus markers were not a significant risk factor implying that, in previous analyses, the risk associated with hepatitis B virus might have been confounded by the coassociation of hepatitis B virus infection with male sex and birth in areas of high incidence. The importance of cirrhosis as a major aetiological factor is now more widely accepted, at least in low hepatocellular carcinoma incidence areas, but the interactions with other aetiological factors including hepatitis B virus infection remain controversial.

SEX HORMONES AND LIVER CANCER

It had been long established that, in certain mouse strains, administration of testosterone significantly increases the frequency with which chemically induced hyperplastic nodules progress to hepatoma and castration of male animals decreases the frequency of this occurrence.8 In 1966 in a preclinical study of steroids to be used in oral contraceptive preparations it was recorded that, ‘... Dr Bosner reported to the Committee (on the Safety of Drugs) that dosage with mestranol was associated with the production of nodules in rats which histologically showed various stages of nodular hyperplasia and the production of hepatoma ...’9 It should therefore perhaps, not have come as a surprise when Baum10 and her colleagues described the development of hepatic adenomas in women taking oral contraceptive steroids, an association soon to become well accepted. The possible role of these agents in malignant liver disease was more controversial but an influential paper by James Neuberger in collaboration with Professor Richard Doll at Oxford suggested a small, but significantly increased, risk of hepatocellular
cancer after prolonged usage. Such observations, together with the previously noted predominance of men among cirrhotic subjects developing hepatocellular carcinoma, focused on the attention of many on the Unit to the role of sex hormones in hepatic neoplasia.

Jawed Iqbal and colleagues were the first to show the presence of androgen receptors in human liver tissue (Fig 1) which were identical to androgen receptors found in prostatic tissue. These data have since been confirmed as has the observation that human hepatocellular carcinoma tissue exhibits much higher concentrations of androgen receptors and αt-reduced androgens than normal liver tissue. During these studies a new circulating sex-steroid binding protein, 'fetal steroid binding protein' was discovered in human fetal liver and the serum of hepatocellular carcinoma patients. This protein was later purified and shown to be present in normal serum. While its role is uncertain, its levels in hepatocellular carcinoma patients appear to correspond with response to therapy.

**DIAGNOSIS AND MONITORING RESPONSE TO THERAPY**

For all the work into serological tumour markers, alphafetoprotein, first described in the serum of a human with hepatocellular carcinoma in 1964 is perhaps still the only one which is widely used in clinical practice. In the presence of a mass lesion in a cirrhotic liver a markedly raised level is sufficient to establish the diagnosis of hepatocellular carcinoma and monitoring of changes in serum levels can be invaluable in assessing response to therapy (Fig 2). None-}

theless, some of the initial enthusiasm for using alphafetoprotein to screen high risk populations for early detection of hepatocellular carcinoma has subsided with the recognition that there is some overlap between levels of alphafetoprotein in hepatocellular carcinoma and uncomplicated chronic liver disease, particularly while the tumour is small.

The fibrolamellar variant of hepatocellular carcinoma was described in detail in 1980 by Craig et al and for the first time a primary malignant liver cell tumour with distinctive histological features (deeply eosinophilic cyto-

plasm and pyknotic nuclei interspersed with parallel bundles of collagen resulting in fibrous septa) had a specific clinical counterpart. It occurs during adolescence, in a non-cirrhotic liver and is invariably alphafetoprotein seronegative and unrelated to hepatitis B virus infection. The prognosis is somewhat better than with other histological types although very few patients will survive more than five years from diagnosis. At the suggestion of Iain Murray-Lyon, vitamin B12 binding protein was measured and in a series of 107 patients with hepatocellular carcinoma was shown to be a specific marker for this particular tumour variant.

**RADIOLOGY**

The advent of clinical ultrasonography has had a great impact. It is now the first line investigation for screening high risk patients and monitoring response of liver tumours to therapy. Much of the original impetus came from Japan reflecting the combination of technological expertise and the high frequency of hepatocellular carcinoma and from where the concept of ‘minute’ carcinomas (usually defined as less than 3 cm in diameter), eminently suitable for resection, derived. There is now an extensive literature from the Far East on the role of lipiodol image enhancement in computed tomography diagnosis. The aim of this technique is to opacify the liver on a cellular, rather than a vascular basis. Lipiodol, an oily based contrast medium, is injected into the hepatic artery at the time of arteriography and sequential computed tomography scanning is carried out (Fig 3). Lipiodol is cleared from normal hepatic tissues but is said to accumulate in malignant tumours because of the ‘leaky’ character of neovascular tissue, coupled with the lack of lymphatic clearance from tumour tissue.
TREATMENT AND PROGNOSIS

There is no doubt that treatment has been the most disappointing area, one which has seen many false dawns. Other than those patients in whom the tumour is detected incidentally or presymptomatically as a result of screening, the outlook has probably changed little for any of the primary liver tumours. Indeed the use of coeliac plexus block for pain and biliary stenting to relieve obstructive jaundice, although not affecting survival, have probably done more for improvement of quality of life than any of the other attempts at treatment.

SURGERY AND LIVER TRANSPLANTATION

Surgery remains the only hope of 'cure' in patients with malignant liver tumours. Over the last 25 years there has been a gratifying reduction in the operative mortality and an increase in the number of patients considered 'operable', particularly in the Far East. This reflects a combination of better surgical and anaesthetic techniques, in particular appreciation of the segmental anatomy of the liver. About 40% of hepatocellular carcinoma patients without cirrhosis and in whom complete resection is obtained can now expect to live for five years, but this group represents only a small minority of the whole population of hepatocellular carcinoma patients. In the Far East surgeons are becoming increasingly enthusiastic about resection in the cirrhotic liver with a five year survival rate of up to 50% being quoted but Western experience with the cirrhotic liver has, to date, been less favourable. Bismuth reported that only 35 of 270 cirrhotic patients were operable and of these, only 12 were alive at 16 months.

Similarly, only a small percentage of tumours in the high bile duct or at the bifurcation are amenable to curative surgical resection. Amongst this group, the development of endoscopic stenting has been a major advance and is now the treatment of choice in most instances where it offers excellent palliation. Low malignant bile duct lesions may also be amenable to endoscopic stenting and a consensus is being reached that this is the treatment of choice in the elderly, frail and ill. The stay in hospital may be shorter and the mortality less than for surgery which should probably be reserved for younger patients with a longer life expectancy.

Surgery is also being increasingly used in patients with metastatic colorectal metastases and guidelines as to which patients will benefit most are becoming available. For example, it appears that for a patient with one or two metastases from a Dukes grade B colonic carcinoma and a resection margin of greater than 1 cm, the chances of five year survival reach almost 50%.

Early experience with liver transplantation for hepatocellular carcinoma was disappointing; whilst the patients did uniformly well in the short term, tumour recurrence was the norm, presumably reflecting spread of clinically undetectable extrahepatic micrometastases to involve the new liver. Our more recent experience, however, has shown that such disappointing results are associated with large tumours and...
that patients with small (<4 cm in diameter) and/or asymptomatic tumours, do much better and seem to be one of the most suitable groups of patients for liver transplantation.

CYTOTOXIC AND ANTI-HORMONAL CHEMOTHERAPY

Early studies on cytotoxic chemotherapy saw the first collaborative ventures between the Royal Free, Charing Cross Hospital, and King's. The first suggested a modest response rate of around 25% with systemic arterial adriamycin in a series of 43 patients. Subsequent studies have not always been able to achieve even such modest levels reported but no other single drug or combination of drugs seem to have fared any better. Recent studies have shown that it is feasible to give full doses of adriamycin even in the presence of hyperbilirubinaemia but the response rate is still disappointingly low.

Our current base line treatment for inoperable hepatocellular carcinoma, against which new therapies should be compared, is to administer three courses of adriamycin systemically at a dose of 60 mg/m² (provided the serum bilirubin concentration is below 30 μmol/l) at three weekly intervals. Where there is objective evidence of response in terms of a decrease in tumour or liver size, or a fall in the serum concentration of alphafetoprotein, the course should be completed (to a maximum dose of 350 mg/m²) otherwise treatment should be withdrawn.

After the detection of sex hormone receptors in hepatocellular carcinoma a prospective randomised controlled trial comparing tamoxifen plus doxorubicin with doxorubicin alone was undertaken. The first patient to enter the study responded dramatically and died, of an unrelated illness whilst in complete remission, three years after. Over the next two years, however, we saw no further such dramatic responses and indeed, in the final analysis, there was no difference in survival. A much more optimistic result using tamoxifen alone has recently been reported with a survival at one year of 40% in the tamoxifen treated group compared with zero in the untreated control group. If these results are confirmed they represent an exciting advance. We have also tried the alternative approach of using the antiandrogen cyproterone acetate in 25 cases. In all five patients with objective evidence of a response there was an associated fall in free 5 α-dihydroprostosterone.

SECONDARY TUMOURS AND THE INFUSEAID PUMP

Several early studies reported that intraarterial chemotheraphy with FUDR led to high remission rates in patients with metastases from colorectal carcinoma but against this needed to be counted the side effects of repeated vascular access. With the development of permanently implantable systems such as the Infuseaid pump these problems seem to have been overcome and recent controlled studies confirm that high remission rates are obtainable. Nonetheless it remains to be proved that these good remission rates are translatable into improved survival, particularly because of the progression of extrahaepatic metastases.

CARCINOID TUMOURS

This period has seen major improvement in the symptomatic control of carcinoid tumours. Although cytotoxic chemotherapy with streptozotocin and 5-fluorouracil is more effective than in hepatocellular carcinoma, it has not been shown to improve survival. As noted above, however, arterial embolisation often gives dramatic symptomatic improvement. Synthetic somatostatin analogues have also led to dramatic improvement in symptoms with less immediate complications than embolisation.

PROGNOSIS

Survival in patients with malignant liver disease is largely dependent on the size of the tumour and the amount of hepatic reserve. The simplest staging classification for hepatocellular carcinoma is that of Okuda (Fig 4a). The prolonged survival of patients with stage I disease should be noted and indeed this is increasing as tumours are being detected earlier by ultrasound or alphafetoprotein screening. It should also be noted that the survival of Japanese patients with stage III disease is similar to that reported from Africa (Fig 4b). This suggests that, rather than having a particularly malignant form of the disease as is sometimes suggested, African patients with hepatocellular carcinoma probably present, or are detected, later than in other countries.

Current areas of interest

The central theme of much of the cancer research at the Institute has been in the still controversial area of the relationship between hepatocellular
carcinoma and cirrhosis. Our prospective follow up study of patients with cirrhosis now involves over 1,400 patients followed for up to 12 years. To date 608 have died, 76 with hepato-cellular carcinoma. This unique database may answer some fundamental questions. For example: suppose the signal for progression to hepatocellular carcinoma occurred simultaneously with that for the development of cirrhosis? This might suggest that even antiviral therapy for hepatitis B virus infection prevented progression to cirrhosis it might still be ineffective in avoiding hepatocellular carcinoma development.

ROLE OF THE HEPATITIS C VIRUS
After identification and cloning of the hepatitis C virus which is responsible for most parenterally transmitted non-A, non-B hepatitis an assay for anti-hepatitis C virus antibodies was developed.48 Up to 80% of patients with hepatocellular carcinoma were reported as seropositive in studies from Italy and Spain together with other types of cirrhosis previously categorised as alcoholic, hepatitis B virus related and autoimmune.49 Ian McFarlane and his co-workers at the United Kingdom have shown that the anti-hepatitis C virus antibody is directed against the hepatitis C virus structural protein and gives results that are comparable to the antibody responses seen in other cirrhosis patients.50-53 Although these results are preliminary and await confirmation by other laboratories, they suggest that determination of a specific antibody response to hepatitis C virus antigens may be of diagnostic help in patients with cirrhosis.

We have recently shown similar results among patients with hepatitis C virus and had the opportunity to test a new diagnostic system based on synthetic peptides derived from both the structural and non-structural part of the hepatitis C virus genome, and this also suggests that only between one and two thirds of the original positive tests can be confirmed. Although these more specific tests have indeed decreased the number of hepatocellular carcinoma patients testing positive for hepatitis C virus on the continent of Europe, a substantial proportion remain hepatitis C virus positive by all available tests. In the United Kingdom it seems likely that its most important implication will be as a cause of a substantial number of cases of 'alcoholic cirrhosis' rather than as a major aetiological factor in hepatocellular carcinoma.

MORE SENSITIVE TUMOUR MARKERS
One of the major potential roles of serum AFP measurement, distinct from its use for diagnosis in symptomatic patients, lies in screening asymptomatic cirrhotic patients with a view to early diagnosis which, as noted above, does seem worthwhile when skillful surgery and/or transplantation are available. A problem soon became apparent, however, small, asymptomatic, tumours tended to have only mildly raised concentrations [in the range 20-500 ng/ml], at which there is some overlap with uncomplexed chronic liver disease. Several studies showed that alpha-fetoprotein derived from patients with chronic liver disease (or other non-malignant conditions such as fulminant hepatic failure or pregnancy) was qualitatively different from that derived from hepatocellular carcinoma in terms of binding to various lectins, particularly lentil lectin. This implied increased fucosylation of the protein and this has been confirmed.54,55 Du and Ming Qiong have recently shown that such abnormalities could be detected even amongst hepatocellular carcinoma patients who had concentrations of alpha-fetoprotein below 400 ng/ml—i.e. that is, concentrations which are, quantitatively not considered diagnostic.56 They also showed that the underlying mechanism of these changes is probably related to disturbances of enzymatic control particularly involving fucose transferases which are very active in both the serum and tissue of hepatocellular carcinoma patients.57

TARGETING OF CYTOTOXIC DRUGS
In attempts to increase selectivity, cytotoxic agents have been linked to polyclonal and monoclonal antibodies and lipiodol, an agent which, as previously noted, localises in tumour tissues after arterial administration. Several groups from Japan and the Far East have suggested that this is an ideal method for targeting cytotoxic drugs, because it leads to increased concentration of drug in the tumour and therefore to increased clinical efficacy and might be expected to decrease systemic toxicity. In an extensive series of studies, Cem Kalayci showed that the co-administration of lipiodol had very little effect on the pharmacokinetics, toxicity or clinical efficacy of adriamycin (Fig 5).57,58 It was concluded that previous favourable results were the result of other methods of treatment administered at the same time as the lipiodol chemotherapy, particularly arterial embolisation.

GROWTH FACTORS
The autocrine mechanism of cellular transformation is probably one of the most frequent initiators of tumourigenesis.59 Cells that possess specific receptors for the polypeptides they secrete, produce either or both ligand and receptor in an uncontrolled fashion. Increased insulin like growth factor II mRNA transcripts have been reported in the hepatocellular carcinoma developing in woodchucks infected with hepatitis B virus and recently, using in situ hybridisation Carolyn d'Arville and Kayhan Nouri-Aria have shown that the expression of insulin like growth factor II mRNA is upreg-
ulated in hepatocellular carcinoma tissue where there is cirrhosis and when the patients are carriers of the hepatitis B virus.65

The future
Assuming that current vaccines remain effective, the eradication of hepatitis B (and later, perhaps hepatitis C) could probably eliminate most hepatocellular carcinoma in the East within 50 years. In the West, control of excessive alcohol consumption would probably be almost as effective. To this extent medical science has offered the solutions—their implementation is dependent on financial considerations and political will.

The search for the mechanism by which hepatitis B virus is oncogenic (if indeed it is!) is the next great prize—there may well turn out to be several different mechanisms. At present it seems most likely that a cellular oncogene is activated by integration of viral DNA and its promoter sequence upstream from the oncogene. Alternatively, viral DNA integration is an essential step but it is subsequently eliminated from the site of action leaving a permanent deletion—the so called ‘hit and run’ hypothesis.

The concept of early diagnosis will perhaps need to be redefined with respect to hepatocellular carcinoma. The results of ‘too late’ diagnosis are only too well recognised; those of ‘too early’ diagnosis are only just being recognised. There is probably just a brief window of opportunity when the tumour is sufficiently large that we can be sure that it poses a significant threat to its host and yet is small enough to be successfully resected. An important recent study suggests that adenomatous hyperplasia may be the true precancerous lesion but detection of such foci will provide major therapeutic dilemmas.64 Hilar cholangiocarcinoma remains a major problem—often impossible to diagnose or treat effectively.

The role of screening high risk populations with a view to early diagnosis remains contentious and its potential value in different countries is being actively investigated. In the United Kingdom it seems unlikely to have a major impact, as the high risk groups (namely carriers of the hepatitis B virus and patients with cirrhosis) are seldom identified before the development of tumour.65

Conclusions
Many of the advances in the field of hepatic oncology have arisen from chance observations in unrelated fields and progress seems to have been haphazard. More ‘goal directed’ research, exemplified by the cloning of the hepatitis C virus by pharmaceutical and pharmaceutical companies can be seen as a way of overcoming the prodigious expense of such research. The downside, however, including reluctance to publish in conventional scientific journals, and the patenting of genetic sequences has become all too apparent. There is no such thing as a free diagnostic test or vaccine; the health service will ultimately have to cover research costs.

If the potential for research in this important and intellectually intriguing area is great, many of those committed to research in a hospital environment have not been reassured by recent Government attitudes which appear to undervalue the role of research as an integral part of the health service. So much of what has been successful in the Institute’s contributions has resulted from wide and free referral of patients even when, as is the case with hepatocellular carcinoma, no definitive therapy is available. The likely deleterious impact of an internal market for patients on such free referral should not be underestimated.

Finally, it is sobering to recall that behind all the Institute’s academic endeavours and publications, each point on the accompanying figures usually represents one person suffering from these devastating forms of cancer. These were all cared for as patients on our wards. The management of those whom we have no way to cure is perhaps the most demanding of all clinical disciplines, and the best reason for continued research.

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