Structures of disease-specific serum alpha-fetoprotein isoforms

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Summary
Alpha-fetoprotein (AFP) is widely used as a serological marker in the diagnosis of hepatocellular carcinoma (HCC) and non-seminomatous germ cell tumours (NSGCT). By application of isoelectric focusing (IEF) disease-specific AFP isoforms can be identified. Three major bands are apparent: + l (associated with ‘benign’ liver disease), + II (associated with HCC) and +III (associated with NSGCT).

Recently, we have characterized the predominant glycans of human serum AFP and now report the application of these findings and electrospray ionization-mass spectrometry (ESI-MS) to the determination of the glycan composition of the isoforms present in the sera of 12 patients with HCC and of one patient with NSGCT. ESI-MS allowed simultaneous identification of various AFP glycoforms in purified serum AFP. Seven glycoforms were identified, but with different abundance in the sera of the HCC patients, whereas six glycoforms were identified in the sera from the NSGCT patient. The glycan structures of these glycoforms were deduced from their observed masses. AFP glycoforms carrying a single biantennary complex-type N-glycan appeared as the predominant glycoforms, whereas those carrying both N-glycan and O-glycan appeared as minor glycoforms. Correlation between the abundance of the AFP glycoforms and the IEF band intensity suggested that different degrees in sialylation cause the formation of isoforms. This contention was subsequently supported by the ESI-MS and kinetic in vitro desialylation studies on purified Bands + I and + II AFPs. Our findings indicate that HCC-associated isoforms (Band + II) represent a group of glycoproteins whose carbohydrate structures are all characterized by being mono-sialylated, whereas those associated with benign liver disease and NSGCT are di- and a-sialo species, respectively. Knowledge of the structure of the tumour-specific isoforms should form an important basis for clinically useful assays. © 2000 Cancer Research Campaign

Keywords: hepatocellular carcinoma; non-seminomatous germ cell tumours; alpha-fetoprotein isoform; protein structure; glycosylation

Measurement of serum alpha-fetoprotein (AFP, reference range < 10 ng ml⁻¹) provides a marker for the diagnosis and management of hepatocellular carcinoma (HCC) (Johnson et al, 1978; Nomura et al, 1985; Sheu et al, 1985) and non-seminomatous germ cell tumours (NSGCT) (Lange and Fralay, 1977; Javadpour 1980; Kay et al, 1987; Nichols et al, 1990). About 70% of patients with HCC will have levels above the reference range (Johnson et al, 1978; Nomura et al, 1985; Sheu et al, 1985). A serum concentration of greater than 500 ng ml⁻¹, in an HCC high-incidence area, and in the appropriate clinical setting, is usually diagnostic of HCC. However, modestly raised levels of AFP (10–500 ng ml⁻¹) are also common in non-malignant chronic liver disease, so that the specificity of the AFP test for HCC tends to be low (Johnson et al, 1978; Okuda, 1986; Lok and Lai, 1989). This represents a serious clinical drawback for the test since most cases of HCC arise in patients with concurrent chronic liver disease (Kew and Popper, 1984; Johnson and Williams, 1987).

AFP is a glycoprotein consisting of 591 amino acids that has been reported to have a single asparagine linked complex-type sugar chain (Tarelli et al, 1992; Ferranti et al, 1995). Although it has been known for over 20 years that serum AFP exhibits microheterogeneity when examined by isoelectric focusing (IEF), the biomolecular basis for this heterogeneity has remained controversial (Ruoslahti 1979; Smith and Kelleher, 1980). Recently, we have used fluorescence labeling, sequential exoglycosidase digestion, high-performance liquid chromatography and matrix-assisted laser desorption ionization in time-of-flight mass spectrometry, to determine the glycan structures of purified serum AFP from patients with HCC and NSGCT. Surprisingly, eleven major glycans were found, of which seven were N-linked, and four were O-linked, to the protein backbone (Figure 1) (Johnson et al, 1999).

Several attempts have been made to identify an ‘HCC-specific’ glycoform, aiming thereby to improve the specificity of AFP as a diagnostic test for HCC. The most successful approach has been based on the difference in the binding affinity of the AFP glycoforms to various lectins, particularly lentil lectin and Concanavalin A. Indeed, different binding patterns were found in AFP from patients with different diseases (Smith and Kelleher, 1980; Buamah et al, 1986; Du et al, 1991). More recently, using IEF to directly identify isoforms of AFP, the Band + II and Band + III AFP isoforms were shown to be highly specific for HCC (Burditt et al, 1994; Ho et al, 1996) and NSGCT (Johnson et al, 1995), respectively. A unique IEF pattern is also found for the AFP that arises in patients with chronic liver disease without any evidence of malignancy (Ho et al, 1996; Johnson et al, 1997). Preliminary studies have suggested that screening for the Band + II isoform may allow early, even preclinical, diagnosis of HCC in high-risk patients (Johnson et al, 1997). However, the current protocol of IEF and immunoblotting techniques for identification of the tumour-specific isoform is very time-consuming and it is therefore
important to develop a simpler and more rapid assay. To achieve this, it is essential to determine the biomolecular basis of the disease-specific isoforms so as to identify any distinctive structural features.

Although the carbohydrate structures of serum disease-specific glycoforms have been studied using the lectin affinity electrophoretic technique (Shimizu et al, 1996), the structures of the HCC and NSGCT isoforms identified by IEF remain unknown. In this study, using electrospray ionization-mass spectrometry (ESI-MS) and applying our recent characterization of the 11 glycans of AFP, we have elucidated the structures of the glycan moieties of those AFP isoforms associated with disease-specific states, and related these to the IEF banding patterns. ESI-MS and kinetic in vitro desialylation studies were then applied to confirm the structures of disease-specific isoforms.

MATERIALS AND METHODS

Patient sera
Sera were collected from 12 patients with histologically confirmed HCC, all of whom had serum AFP levels of > 300 000 ng ml⁻¹ (range 300 000–300 000 ng ml⁻¹), and one patient with histologically confirmed NSGCT with a serum AFP level of 17 820 ng ml⁻¹. The concentration of serum AFP was assayed by a microparticle enzyme immunoassay (MEIA, Abbott Laboratories, Illinois, USA). Sera were stored at −70°C until further analysis, or until purification of the AFP.

Purification of AFP by one-step affinity chromatography

Rabbit anti-human AFP (DAKO, Glostrup, Denmark) was coupled to CNBr-activated Sepharose 4B (Pharmacia Biotech, Uppsala, Sweden). A mini-column containing 1 ml of coupled gel was equilibrated with phosphate buffered saline (PBS, 10 mM sodium phosphate, 10 mM KCl, 127 mM NaCl, pH 7.4). The sample to be purified (200–500 μg) was loaded into the column and incubated at room temperature for 4 h. The column was then washed with 50 ml of PBS containing 0.2% Tween 20, and subsequently with 50 ml of PBS. Bound AFP was eluted with 20 ml of 0.2 M Na₂CO₃ under gravity. The ‘total’ AFP concentration was determined as previously described. The purity of the final AFP preparation was regarded as acceptable when a single band, with a molecular weight of about 68 kDa, was visualized on an SDS PAGE gel using silver staining.
Purification of AFP isoforms by preparative IEF and affinity chromatography

Serum containing about 5 mg of AFP was applied to a mixture of 5% sorbitol, 10% glycerol and ampholytes (6.25% Pharmalyte pH 4.5–5.4, Pharmacia Biotech) which had been pre-focused at constant power (15 W) for 30 min in a Rotofor cell (Biorad, California, USA) at 4°C. The sample was then focused at constant power (15 W) until the voltage stabilized and then focused for a further 30 min before 20 individual 2 ml fractions were harvested. pH and AFP concentration in each fraction were measured. Fractions with more than 10 μg of AFP were pooled and refractionated to improve the separation of the isoforms. The pH and AFP concentration were again measured. Fractions containing only Band +I or Band +II AFP, free of other isoforms, were identified by gel IEF. Finally, the separated AFP isoforms in the pooled fractions were purified by affinity chromatography.

Isoelectric focusing-immunoblotting (IEF-IB)

The method of Burditt et al (1994) with minor modifications was used. Samples were focused in 1.5 mm thick agarose gel containing 1% agarose (IEF grade Type VIII, Sigma, Missouri, USA) 5% sorbitol, 10% glycerol and ampholytes (6.25% Pharmalyte pH 4.5–5.4, Pharmacia Biotech) at 10°C. After pre-focusing at 1500 V for 30 min, samples were then applied directly to the gel and focused at 2000 V for 1 h. The focused proteins were transferred to nitrocellulose membrane, and then incubated with horseradish peroxidase conjugated polyclonal rabbit anti-human AFP (DAKO). After washing, the enhanced chemiluminescence detection system (ECL, Pharmacia Biotech) was used to visualize the AFP protein bands.

ESI-MS analysis of purified AFPs

ESI-MS was performed on a triple quadrupole Quattro II system (Micromass, Manchester, UK). The mass scale was calibrated with a mixture of myoglobin and trypsinogen. The capillary voltage of the electrospray probe was set to + 3.5 KV. The nebulizer gas was set at 20 l h–1 and the drying gas at 250 l h–1. The source temperature was set at 75°C. The desalted AFP sample (5 pmol μl–1 in 50% acetonitrile in water containing 0.2% formic acid) was injected into the ion source at a flow rate of 5 μl min–1. Positive ionization mass spectra were acquired over the mass/charge ratio (M/Z) range of 1000–2400 at 4 s per scan. The sampling cone voltage was varied through the range of 47–90 V. The background subtracted raw data was processed by the Maximum Entropy (MaxEnt) software to produce the transformed mass spectrum.
The accuracy of this method is $\pm 0.01\%$. All samples were measured for 3–5 times.

Deduction of the glycan composition in AFP glycoforms identified by ESI-MS

With the knowledge of the structures of eleven predominant glycans on AFP glycoforms, the molecular masses for a series of hypothetical AFP glycoforms comprising random combinations of the $N$-linked and/or $O$-linked glycans were calculated. Their theoretical masses (average masses) were derived from the theoretical mass of the AFP protein core (calculated with the Peptide Mass software from ExPASy, accessed through the Internet at http://expasy.hcuge.ch), and the theoretical masses of the monosaccharide residues reported by Dell et al (1993). The observed masses of the AFP glycoforms identified by ESI-MS were then matched with the calculated masses of the hypothetical AFP glycoforms. Those observed AFP glycoforms in which mass difference was less than $0.01\%$ were considered to have the same molecular structure as the matched hypothetical AFP glycoforms.

The accuracy of this method is $\pm 0.01\%$. All samples were measured for 3–5 times.

### Table 2

| Observed mass | Deduced glycan composition | Calculated mass | Monosaccharide composition per protein molecule | Label |
|---------------|---------------------------|----------------|-----------------------------------------------|-------|
| 68803 ± 1.5   | G7                        | 68800.5        | 2 SA, 5 Hex, 4 HexNAc, 1 DeoxyHex              | +Ib   |
| 68712 ± 0.8   | G10 + G2b + G2b           | 68712.0        | 1 SA, 5 Hex, 5 HexNAc, 1 DeoxyHex              | +IIb  |
| 68511 ± 1.6   | G8                        | 68508.8        | 1 SA, 5 Hex, 4 HexNAc, 1 DeoxyHex              | +Ic   |
| 68342 ± 2.3   | G11 + G2b                 | 68346.7        | 1 SA, 4 Hex, 5 HexNAc, 1 DeoxyHex              | +Ie   |
| 68423 ± 0.6   | G10 + G4                  | 68420.7        | 5 Hex, 5 HexNAc, 1 DeoxyHex                    | +IIla |
| 68053 ± 1.1   | G10                       | 68055.4        | 4 Hex, 4 HexNAc, 1 DeoxyHex                    | +IIb  |

*First predominant glycoform; second predominant glycoform; third predominant glycoform; most probable glycan composition within the group

### Table 3

| HCC case | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
|----------|---|---|---|---|---|---|---|---|---|----|----|----|
| AFP      | +Ia | +Ia | +Ib | +Ib | +Ib | +Ib | +Ib | +Ia | +Ia | +Ia | +Ia | +Ia |
| Glycoforms | +Ic | +Ic | +Ic | +Ic | +Ic | +Ic | +Ic | +Ic | +Ic | +Ic | +Ic | +Ic |
|          | +Ib | +Ib | +Ib | +Ib | +Ib | +Ib | +Ib | +Ib | +Ib | +Ib | +Ib | +Ib |
|          | +IIa | +IIa | +IIa | +IIa | +IIa | +IIa | +IIa | +IIa | +IIa | +IIa | +IIa | +IIa |

*Predominant AFP glycoform; second predominant AFP glycoform

Kinetic in vitro desialylation of purified Band + I AFP

20 µl of 250 mU ml$^{-1}$ sialidase (Vibrio cholerae neuraminidase, Roche Diagnostics, Indianapolis, USA) was mixed with 10 µl of purified Band + I AFP (800 µg ml$^{-1}$) in reaction buffer (0.1 M Na acetate, 9 mg ml$^{-1}$ NaCl, 1 mg ml$^{-1}$ CaCl$_2$, pH 5.6), and incubated at 37°C. At the assigned time for the individual assay tubes, the reaction was stopped by adding 20 µl of stopping buffer (0.15 M glycine-NaOH, pH 11.0). The final reaction mixture was subjected to IEF-IB for the visualization of the AFP IEF banding patterns.

### RESULTS

Identification of glycan composition of AFP glycoforms in patient sera by ESI-MS

Total AFP was successfully purified from individual patient sera (Figure 2), and subjected to ESI-MS analysis. Four major and three minor AFP glycoforms, purified from the sera of the HCC patients, were identified (Table 1). Three major and three minor
AFP glycoforms were identified in the serum from the NSGCT patient (Table 2). Their compositions were deduced from the observed molecular masses after being matched with the hypothetical AFP glycoforms as described above. In total, 10 AFP glycoforms were identified from the patient sera. For seven out of the 10 identified AFP glycoforms, there was only one possible glycan composition that could be deduced from the observed mass for each of the glycoforms.

Although three out of 10 identified AFP glycoforms (AFPs + Ib, + Ib and + IId) were matched with more than one hypothetical AFP glycoforms (Tables 1 and 2), the comparison with those identified glycoforms with concrete glycan moiety(s) can provide a clue to decide which one is the most probable glycoform. In the case of AFP + Ib, its expected theoretical mass is greater than that of AFP + Ic by 146.1, which is equivalent to the mass of a fucose. It is possible that AFP + Ib has a glycan moiety similar to AFP + Ic, but with an additional fucose. Such postulation helps to explain why only either AFP + Ib (67%, 8 of 12) or AFP + Ic (33%, 4 of 12) appeared as the predominant glycoforms in sera from HCC patients (Table 3). The presence of AFP + Ib as the predominant glycoform may be caused by a higher fucosylation activity in hepatoma cells. Therefore, the fucosylated disialo biantennary complex-type N-linked glycan (G7) is the most probable glycan moiety of the AFP + Ib, which is consistent with the identification of G7 glycans as one of the major glycans released from HCC-associated AFP (Yoshima et al, 1980; Aoyagi et al, 1993; Johnson et al, 1999). In the case of AFP + IId, it was only found in sera with a negligible amount of AFP + Ic (Table 3). The expected theoretical mass of AFP + IId is greater than that of AFP + Ic by 203.2, which is equivalent to the mass of an HexNAc residue (G5). The absence of the AFP + Ic could be possibly explained by the addition of a G5 glycan to AFP + IId, converting it to AFP + IId. Therefore the most probable glycan moieties of AFP + IId are G8 and G5 glycans. For the AFP glycoforms identified in the NSGCT serum, because G10 glycan is the sole contributor of the carbohydrate content of the second predominant glycoform, + IId, it is reasonable to postulate that the AFP + IId also carried one G10 glycan, but with an additional G4 glycan (Table 2). Thus, it appears that glycoforms carrying a G10 glycan predominated in the NSGCT-derived AFP, and contributed to glycoforms + IId, + IId and + IId.

Classification of the identified AFP glycoforms

Since it is known that the isoelectric point (pI value) a glycoprotein is affected by the number of sialic acid residues (Stibler, 1991), the identified AFP glycoforms were classified according to their sialic acid content – Class +I, Class +II and Class +III, carrying two, one and no sialic acid residues, respectively. In Class +I there were three subtypes (+ Ia–Ic); in Class +II there were five subtypes (+ IIa–IIe); in Class +III there were two subtypes (+ IIIa and + IIIb). The classification of the identified AFP glycoforms from sera of the HCC patients and from the NSGCT patient are summarized in

Table 4  The major serum AFP glycoforms, identified by ESI-MS, in AFP preparations that were purified from two HCC patients. Total AFP, Band +I AFP and Band +II AFP were purified by affinity chromatography and preparative IEF, and subsequently subjected to ESI-MS analysis for 3–5 determinations. The glycan compositions of the identified glycoforms were deduced from their observed masses.

| Patient | Total AFP | Band +I AFP | Band +II AFP | Observed mass by ESI-MS | Identity | Deduced glycan composition |
|---------|-----------|-------------|--------------|------------------------|----------|--------------------------|
| Patient 1 | 68361 | – | 68362 | AFP +IId (68362.7) | G6 |
| 68655 | 68648 | – | | AFP +Ic (68653.9) | G3 |
| 68801 | 68797 | – | | AFP +Ib (68800.0) | G7 |
| 68352 | – | 68355 | | AFP +IId (68362.7) | G6 |
| 68507 | – | 68505 | | AFP +Ic (68506.3) | G8 |
| Patient 2 | 68651 | 68655 | – | | G3 |
| 68797 | 68802 | – | | | G7 |

*a appearing as a minor glycoform
Tables 1 and 2, respectively. After grouping according to the sialic acid content, it was apparent that all the identified glycoforms had similar carbohydrate contents (9–11 neutral monosaccharide residues). The similarity in the protein core and amount of neutral monosaccharides in all the identified glycoforms strongly suggest that differences in the pl values of the glycoforms are dictated mainly by the number of sialic acid residues.

**Distribution and abundance of the identified AFP glycoforms in different patients**

The distribution of the AFP glycoforms in the sera of patients with HCC is presented in Table 3. The pattern is heterogeneous but several consistent features emerge. In all cases Class + I AFPs predominated, whereas Class + II AFPs appeared as the second predominant glycoforms. AFP glycoforms carrying a single disialylated biantennary complex-type N-glycan (G3 or G7) predominated in Class + I AFPs, whereas AFP glycoforms carrying a single monosialylated biantennary complex-type N-glycan (G6 or G8) predominated in Class + II AFPs. AFP glycoforms carrying both the N-glycan and O-glycan appeared as minor glycoforms in the patient sera.

Owing to the rarity of NSGCT patient serum samples with sufficiently high levels of AFP for analysis, we only have been able to use ESI-MS to elucidate the structure of AFP glycoforms derived from a single case of NSGCT. No claim, therefore, has been made for the predominant glycoforms of NSGCT-associated AFP. However, AFP glycoforms carrying an asialylated biantennary complex-type N-glycan (G10) appeared as the predominant glycoforms in Class + III AFPs.

**Deduction of glycan composition of AFP isoforms identified by IEF**

In all serum samples from the HCC patients, comparison of the intensities of Band + I and Band + II showed that Band + I predominated (Figure 2). Similarly, ESI-MS identified Class + I AFPs as the predominant glycoforms (Table 3). By matching the abundance, we postulate that Class + I and Class + II AFPs represent Band + I and Band + II on the IEF gel, respectively (Figure 2 and Table 3). Similarly in the case of NSGCT, Class + II and + III AFPs should represent Bands + II and + III, respectively (Figure 2 and Table 2).

As the addition of one sialic acid residue decreases the pl value (of a glycoprotein) by about 0.1 pH (Stibler, 1991), Class + I AFPs should, if our postulation is correct, have a pl value lower than that of Class + II AFPs by about 0.1 pH, and the pl value of Class + II AFPs lower than that of Class + III AFPs, again by about 0.1 pH. Consistent with this prediction, the pl values of the IEF bands were: Band + I = pl 4.78, Band + II = pl 4.83 and Band + III = pl 4.90 (Figure 2). The proposed distribution of Class + I, Class + II and Class + III AFPs on the gel IEF pattern is summarized in Figure 3. It is apparent that more than one subtype of each class of AFPs usually coexisted in individual patient serum (Tables 2 and 3). Furthermore, thick bands were usually seen on the IEF gel (Figure 2). We suggest that, as summarized in Figure 3, several AFP isoforms, all with very close pl values (those within a class), are focused at similar positions. Their bands overlap with each other on the IEF gel, and finally appear as a single thick band.

**Identification of glycan compositions of purified Band + I and Band + II AFPs by ESI-MS**

To further support our postulated structures for the specific IEF bands, besides the total AFP we prepared purified Band + I and + II AFPs from two HCC patients (Figure 2), and subjected them to ESI-MS analysis. Unfortunately, the level of Band + III AFP in the NSGCT serum was too low to allow us to purify a sufficient
amount for ESI-MS analysis. By ESI-MS (Figure 4 and Table 4), both Class +I (putatively disialylated) and Class +II (putatively monosialylated) AFS were identified in the purified total AFP preparations (comprising all the AFP glycoforms existing in the sera). As predicted, Class +I (disialylated) AFS were, indeed, predominantly retained in the Band +I AFP preparations, whereas the Class +II (monosialylated) AFS were predominantly retained in the Band +II AFP preparations. However, fucosylated and non-fucosylated AFP glycoforms were not differentially separated by preparative IEF. The results of these ESI-MS experiments indicate that Class +I (disialylated) and Class +II (monosialylated) AFS were focused as Band +I and +II AFS, respectively.

**Kinetic in vitro desialylation of purified Band +I AFP**

In the presence of sialidase, the purified Band +I AFP, in a step-wise manner, migrated from the position +I to +II and finally to +III (Figure 5). Consistent findings were observed on two independent occasions. Upon partial sialidase digestion (0.5 h of digestion), microheterogeneity increased immediately, showing a new alkaline band at position +II. After 2.5 h of digestion, most of Band +I AFP was converted to the glycoform at position +II. After 5 h of digestion, another glycoform at position +III appeared. Upon complete digestion (20 h of digestion), the majority of the AFP protein molecules (asialylated AFS) were still focused as Band +III. Non-specific effects of incubation conditions were excluded by control experiments. Conversion of Band +I AFP to Band +II and +III AFP upon sequential removal of sialic acid residues confirmed that the appearance of alkaline AFP glycoforms (+II and +III) identified by IEF were attributable to a lower sialic acid content, and asialylated AFP was focused as Band +III. Thus, this kinetic in vitro desialylation experiment strongly supports our contention that Band +I, +II and +III are composed of di-, mono- and a-sialylated AFS, respectively.

**DISCUSSION**

Although AFP has been used as a tumour marker for the diagnosis of HCC for many years, its specificity is poor when levels fall within the ‘grey area’, i.e. 10–500 ng ml⁻¹ (Johnson et al, 1978; Okuda, 1986; Lok and Lai, 1989). In this range, differentiation between benign and malignant liver diseases cannot be made with confidence on the basis of serum AFP levels alone. There is thus a need to find a method of increasing the specificity of AFP. Lentil lectin-reactive AFP has previously been proposed as a diagnostic marker to discriminate between AFP arising from HCC and AFP arising from benign liver disease (Du et al, 1991; Taketa et al, 1993; Shimizu et al, 1996). Using the lectin binding properties of N-glycans with known structures (Yamashita et al, 1993) or glycodiesease digestion (Shimizu et al, 1996) together with lectin affinity electrophoresis, the carbohydrate structure of the major HCC-specific AFP glycoform has been suggested indirectly to be of the fucosylated monosialo biantennary complex-type.

Identification of AFP isoforms is an alternative method to increase the specificity of AFP. Using IEF and immunoblotting, AFP isoforms appearing as Band +I and Band +III were found to be relatively specific for HCC and NSGCT, respectively (Johnson et al, 1995; Ho et al, 1996; Johnson et al, 1997). The direct structural analysis of these AFP isoforms in individual patients has, until recently, been hindered by the extremely small amounts of serum AFP available. However, with the help of ESI-MS and a random assembly model (i.e. the construction of hypothetical AFP glycoforms from the AFP protein core and the known glycans identified on AFP), we report here an approach that has permitted simultaneous identification of the predominant AFP glycoforms in individual patients. The present study provides evidence that the glycosylation of human serum AFP is more complex than the single biantennary complex-type N-glycan that has previously been reported. There exist AFP glycoforms carrying both O-glycan and N-glycan. The identification of AFP glycoforms carrying O-glycan was consistent with our previous identification of O-glycans released from AFP molecules (Johnson et al, 1999).

Our results show that an AFP glycoform carrying a single monosialo biantennary complex-type N-glycan, either fucosylated or not, is focused as Band +II. As Band +II AFP is, on the basis of our previous studies, specific for HCC, our findings suggest that aside from fucosylated monosialo biantennary complex-type N-glycan (Yamashita et al, 1993; Shimizu et al, 1996), non-fucosylated monosialo biantennary complex-type N-glycan is also the predominant glycan moiety of the HCC-specific AFP glycoforms. The AFP glycoform carrying a single difucosylated disialo biantennary complex-type N-glycan, which had been identified in a hepatoblastoma cell culture (HepG2) by ESI-MS (Ferranti et al, 1995), was not found in our patient sera. Further investigation is necessary before we can answer whether hyper-fucosylation of AFP is a unique property of hepatoblastoma, but not to HCC, or whether it is due to change in cellular properties upon in vitro cell culture.

Among the predominant AFP isoforms, in our IEF system, the one that is commonly identified together with Band +II AFP in sera of HCC patients and usually occurs alone in sera of patients with benign liver disease is Band +I AFP (Ho et al, 1996; Johnson et al, 1997). This ‘benign’ isoform appears to be composed of one disialylated biantennary complex-type N-glycan. As fucosylation of the N-linked glycan has been postulated to happen only during oncogenesis because of loss of cell polarity (Yamashita et al, 1993), the non-fucosylated disialo biantennary complex-type N-glycan is likely to be the main carbohydrate structure of this ‘benign’ AFP. Unfortunately, the amount of serum AFP in patients with liver cirrhosis is always too low (< 500 ng ml⁻¹) to permit purification for the purpose of structural analysis. With the currently available technology it is therefore difficult to confirm our suggested structure for the ‘benign’ AFP isoform.

It has been reported that detection of lentil lectin-reactive AFP by lectin affinity electrophoresis may only discriminate benign liver diseases from HCC with a maximum diameter of nodules > 20 mm, but not from early malignancy (maximum diameter of nodules <10 mm) (Taketa et al, 1993). In contrast, our preliminary studies have suggested that detection of Band +II AFP may allow early diagnosis of HCC in high-risk patients even before it can be detected by routine ultrasound examination (Johnson et al, 1997). The detection of lentil lectin-reactive AFP may thus be less sensitive than Band +II AFP for early diagnosis of HCC. The appearance of monosialo-AFP (Band +II AFP) in the blood during the development of HCC in patients with chronic liver disease or cirrhosis may, in part, be related to a decrease in the sialylation activity in HCC cells. As a result, the apparent higher specificity of monosialo-AFP suggests that the decrease in the sialylation activity may occur earlier than the increase in the fucosylation activity during oncogenesis in human liver.

In conclusion, three classes of AFS, +I, +II and +III, are identified in patient sera by ESI-MS. The predominant AFP glycoform
in each class carries one disialo, monosialo, or asialo biantennary complex-type N-glycans, respectively. These three classes correspond to ‘benign AFP’ (appearing as Band +I), ‘HCC-specific AFP’ (Band +II) and ‘NSGCT-specific AFP’ (Band +III), respectively. Knowledge of the structure of the tumour-specific isoforms should form the basis for clinically useful assays.

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