Study of childhood renal tumours using peroxidase conjugated lectins

S KUMAR, T CARR, HB MARSDEN, PH MORRIS-JONES

From the Christie Hospital and Holt Radium Institute, Manchester

SUMMARY Six peroxidase conjugated lectins were used to compare their ability to bind to formalin fixed paraffin embedded tissue sections of childhood renal tumours (Wilms' tumour, mesoblastic nephroma, renal carcinoma, rhabdoid renal tumour, and bone metastasising renal tumour of childhood (BMRTC)) with fetal and normal children's kidney. Lectins were found to be helpful in the differential diagnosis of renal tumours. Another important finding was that the mesenchyme of renal tumours showed differences in its reactivity among various types of kidney tumours. The results of lectin binding were not helpful in establishing the origin of kidney tumours.

Lectins are a group of naturally occurring proteins that bind specifically to carbohydrate components of glycoproteins.1 In human kidney, fluorescence or peroxidase labelled lectins have been shown to be capable of distinguishing various anatomical components of a nephron3–7: winged pea lectin (LTA) binds specifically to cells of proximal tubules, and horse gram lectin (DBA) to those of distal tubules but not vice versa. The histogenesis of children's kidney tumours remains unresolved, or at least is controversial.8–9 Wilms' tumour or nephroblastoma, which is the most common renal neoplasm among children, is considered to possess markers of both proximal and distal tubules. Renal carcinoma is the most common renal tumour in adults but its incidence in children is only about 3% of all renal neoplasms. The origin of this tumour type remains uncertain.8–10–13 Some authors consider that it resembles proximal tubules, whereas others maintain that it shares common features with distal tubules. The histogenesis of three other rare types of children's renal tumours—namely, mesoblastic nephroma, rhabdoid renal tumour, and bone metastasising renal tumour of childhood (BMRTC) has not been accurately ascertained.8 To gain some insight into the development of kidney neoplasms we compared the staining reactivity of these tumours with normal human fetal and children's kidneys, using a panel of six lectins. In addition, the results of lectin staining were also analysed to examine how lectin binding carbohydrates were distributed within one histological tumour type and between different types of childhood renal tumours.

Material and methods

LECTINS Six different peroxidase conjugated lectins were purchased from Sigma. Table 1 lists the lectins used, together with their sugar specificities and concentrations found to be optimal for staining in the present study.

TISSUES Tumours were either fresh biopsy or necropsy specimens obtained within a few hours of death. Likewise, normal kidneys were collected from children either at the time of operation or at necropsy. Fetal kidneys were collected soon after therapeutic abortion (14–21 weeks of gestation). All tissues were fixed in 10% buffered formalin, embedded in paraffin, and sectioned at 5 μm thickness.

LECTIN HISTOCHEMISTRY The peroxidase staining method used was that of Polak and Van Noorden.14 Paraffin embedded sections were dewaxed, and endogenous peroxidase was inhibited using 0.5% hydrogen peroxide in methanol. The sections were thoroughly washed in phosphate buffered saline and treated with peroxidase labelled lectin for 30 minutes at room temperature, The slides were washed in phosphate buffered saline, and the colour was developed using freshly prepared diaminobenzidine (Sigma). Sections were counterstained with haematoxylin, dehydrated, cleared, and

Accepted for publication 5 March 1986
Renal tissues and lectins

Table 1  Some details of six peroxidase conjugated lectins used to stain renal tissue

<table>
<thead>
<tr>
<th>Latin name</th>
<th>Common name (abbreviation)</th>
<th>Sugar specificity</th>
<th>Concentration used (µg/ml)</th>
<th>Inhibitory sugar</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arachis hypogaea</td>
<td>Peanut (PNA)</td>
<td>gal-galNac</td>
<td>50</td>
<td>D-galactose</td>
</tr>
<tr>
<td>Dolichos biflorus agglutinin</td>
<td>Horse gram (DBA)</td>
<td>galNac</td>
<td>50</td>
<td>N-acetylgalactosamine</td>
</tr>
<tr>
<td>Lotus tetragonolobus agglutinin</td>
<td>Asparagus or winged pea (LTA)</td>
<td>α-L-fucose</td>
<td>50</td>
<td>L-fucose</td>
</tr>
<tr>
<td>Triticum vulgare</td>
<td>Wheat germ (WGA)</td>
<td>α/β-D-glNac</td>
<td>25</td>
<td>Diacetylchitobiose</td>
</tr>
<tr>
<td>Canavalia ensiform</td>
<td>Jack bean (CON A)</td>
<td>α-D-mannosyl</td>
<td>10</td>
<td>2-methyl-D-manno-pyranoside</td>
</tr>
<tr>
<td>Ricinus communis type 11-P</td>
<td>Castor bean (RCA)</td>
<td>α/β-D-gal</td>
<td>10</td>
<td>D-galactose</td>
</tr>
</tbody>
</table>

mounted.

To confirm the specificity of lectin binding control sections were treated with the appropriate inhibitory sugar before peroxidase labelled lectins were added (Table 1).

Results

Table 2 gives the results of the lectin staining of fetal and children’s kidney and childhood renal tumours. Normal human fetal and children’s kidneys stained similarly for most of the lectins. A few important differences, however, were observed. Of the six lectins, *Ricinus communis* exhibited the greatest differences in staining. In children’s kidneys *Ricinus communis* stained proximal and distal tubules and some of the collecting ducts. In contrast, *Ricinus communis* staining in fetal kidney was mainly limited to proximal tubules. All cells lining collecting ducts in children’s kidney stained uniformly with horse gram lectin, whereas in fetal kidney every third or fourth cell stained much more intensely than the cells in their neighbourhood. (Fig. 1). Horse gram lectin stained only the distal tubules of children’s kidneys. Winged

<table>
<thead>
<tr>
<th>Lectins used</th>
<th>Staining of normal and fetal kidneys</th>
<th>Staining of children’s kidney tumours</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Kidney source</td>
<td>Tubules</td>
</tr>
<tr>
<td>Peanut (PNA)</td>
<td>Children’s</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>Fetal</td>
<td>–</td>
</tr>
<tr>
<td>Horse gram (DBA)</td>
<td>Children’s</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Fetal</td>
<td>–</td>
</tr>
<tr>
<td>Winged pea (LTA)</td>
<td>Children’s</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>Fetal</td>
<td>+</td>
</tr>
<tr>
<td>Wheat germ (WGA)</td>
<td>Children’s</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>Fetal</td>
<td>+ (Weak)</td>
</tr>
<tr>
<td>Jack bean (CON A)</td>
<td>Children’s</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>Fetal</td>
<td>+</td>
</tr>
<tr>
<td>Castor bean (RCA)</td>
<td>Children’s</td>
<td>+ +</td>
</tr>
<tr>
<td></td>
<td>Fetal</td>
<td>+ +</td>
</tr>
</tbody>
</table>

BMRTC: Bone metastasising renal tumour of childhood; + + strongly positive; + positive; – negative.
Fig. 1  Fetal kidney with horse gram lectin (DBA): note some of cells lining collecting duct are stained much more intensely than cells in their neighbourhood.

Fig. 2  Tubular Wilms' tumour with wheat germ lectin (WGA): tubular lumen is strongly stained.

Fig. 3  Blastemal Wilms' tumour with wheat germ lectin (WGA): strong staining of isolated large cells is apparent in this photomicrograph. Nature of positive cells is not known, but they do not look like infiltrating white cells.

Fig. 4  Renal carcinoma with wheat germ lectin (WGA): strong staining of tumour cells is obvious in this preparation.

Figs. 1-4  Localisation of various peroxidase conjugated lectins in normal children's kidneys, human fetal kidneys, and childhood renal tumours.
pea lectin stained the proximal tubules of both fetal and children's kidneys, but the staining was more pronounced in the proximal tubules of children's kidneys.

In Wilms' tumour peanut lectin, wheat germ lectin, and concanavalin A stained tubule lumens (Fig. 2). The staining reaction seen with horse gram lectin and concanavalin A was interesting: tubular and blastemal cells were negative, whereas the stroma of tubular Wilms' tumours was strongly positive. Winged pea lectin decorated only cystic tubules. Wheat germ lectin not only stained the lumens of tubular Wilms' tumour but also stained isolated large cells in blastemal areas (Fig. 3). The nature of these cells is not known, but they did not look like infiltrating white cells. Ricinus communis stained neither blastemal nor tubular subtypes of Wilms'.

Mesoblastic nephromas were weakly reactive with wheat germ lectin, concanavalin A, and Ricinus communis, whereas horse gram lectin only stained cells in differentiated areas of this tumour type.

Rhabdoid renal tumour stained well with wheat germ lectin, whereas staining was only weak with Ricinus communis and negative with the other lectins.

Renal carcinomas were stained strongly by both concanavalin A and wheat germ lectin but only weakly by horse gram lectin and not at all with the other lectins (Fig. 4).

BMRTC failed to stain by all the six lectins used. Winged pea lectin, however, reacted very strongly with the stroma of this tumour type.

Control tissue sections were not stained when treated with horseradish peroxidase substrate alone (without lectin). Inhibitory sugars abolished staining by the corresponding lectins. Similarly, 1-0% periodic acid treatment inhibited staining by five of the six lectins (wheat germ lectin staining was not inhibited).

Discussion

Several studies have investigated the usefulness of biochemical and immunological markers in human kidney tumours. The existence of antigens associated with Wilms' tumour has been recognised by at least four groups of workers. A Wilms' tumour antiseraum raised by Burtin and Gendron recognised a glycoconjugate. Kumar et al produced an anti-Wilms' tumour antiserum in rabbits that had been rendered tolerant in utero with pooled normal kidney extracts. The antiserum stained fetal kidney, most Wilms' tumours, renal carcinomas, and mesoblastic nephroma but not BMRTC. Wallace and Nairn used two antisera to study renal carcinomas and Wilms' tumours. One of the antisera reacted with brush border protein of proximal tubules and the other, Tamm-Horsfall protein, decorated distal tubules. Although lectins possess a remarkable specificity for binding to sugars even in routinely formalin fixed and paraffin embedded tissues, somewhat surprisingly, no previous study has compared the reactivity of lectins to childhood renal tumours with that to fetal and children's kidneys. From published reports it is also apparent that authors disagree as to whether or not the same lectin stains a particular structure in normal kidney. Halthofer et al found that peanut lectin stained distal tubules of normal kidney, whereas Readler et al found that it did not. As far as we know, no study has been made of fetal human kidney staining by lectins. It is not therefore possible to discuss the present results with reference to any published reports. The results of other relevant studies, however, have been considered where appropriate.

Morphologically, nephroblastoma (Wilms' tumour) is an epithelial and sarcomatous tumour that contains blastemal elements. The proportion of one or the other component can vary greatly between different tumours, or even in the different parts of the same tumour. The presence of tubular differentiation is regarded as a good prognostic feature, whereas a preponderance of blastemal elements indicates a moderately good prognosis. Wilms' tumour is thought to arise from metanephric blastema and has been shown to react with antisera to brush border and Tamm-Horsfall protein antigens, which stain proximal and distal tubules, respectively. In both fetal and children's kidneys several lectins differed as regards their staining of proximal and distal tubules (Table 2). Horse gram lectin, which clearly stained the distal tubules of children's but not fetal kidney, failed to stain either the tubular or blastemal components of Wilms' tumour. The only staining of Wilms' tumour using this lectin was of mesenchyme. Winged pea lectin, on the other hand, reacted only with proximal tubules in children's and fetal kidney but did not stain Wilms' tumour. The distribution of sugars in tubular structures of normal and fetal kidneys and Wilms' tumours seems to differ. This may well be a result of the neoplastic transformation of cells in Wilms' tumours.

Mesoblastic nephroma is a neoplasm of infants, which has an excellent prognosis and requires no chemotherapy or radiotherapy. It comprises 13%-17% of renal tumours in the first year of life. It has been proposed that mesoblastic nephroma is a benign counterpart of a BMRTC and that mesoblastic nephroma arises from secondary rather than primary mesenchyme, which is no longer capable of tubular differentiation. We believe, however, that mesoblastic nephroma originates from primitive nephrogenic mesenchyme before its induction by ure-
teric bud and that it is unlikely to be a precursor of BMRTC. Mesoblastic nephromas were stained by wheat germ lectin, concanavalin A, and Ricinus communis. Their lack of reactivity with horse gram lectin and winged pea lectin (both these lectins react with proximal tubules of normal and fetal children's kidneys) suggests that this tumour probably does not originate from tubular components of the kidney.

Rhabdoid renal tumour is a rare tumour and is usually associated with poor prognosis. The name rhabdoid may well be a misnomer as this tumour type contains two typical epithelial cell markers—epithelial membrane antigen and cytokeratins. We proposed previously that rhabdoid renal tumours arise from committed metanephric blastema. The staining of rhabdoid renal tumour by wheat germ lectin is interesting. Although wheat germ lectin has a wide range of reactivity for various structures in normal fetal and children's kidney, it did not stain tubular or blastemal Wilms' tumour cells (Table 2). The use of this lectin should therefore prove valuable in the differential diagnosis of Wilms' and rhabdoid renal tumour.

BMRTC is mainly a tumour of boys, and features that characterise these neoplasms are the absence of blastema and epithelial structures and the fact that the only mesenchymal differentiation is fibroblastic. Ultrastructurally, BMRTC has been shown to resemble mesangial cells and primitive mesenchymal cells. In vitro it has been shown to possess a strong tendency to invade native collagen substratum. Somewhat surprisingly, all six lectins used failed to decorate cells of BMRTC: the only staining seen was that of mesenchyme by winged pea lectin.

The content of published reports is controversial as regards the origin of renal carcinoma. Some authors consider that it resembles proximal tubules, whereas others note its similarity to distal tubules. Perhaps the situation is more complex than has hitherto been thought. Tannenbaum, from his own studies and those of others, concluded that the granular cell tumour variant of renal carcinoma was closely aligned to proximal tubules and the clear cell variant to distal tubules. In our study all renal carcinomas studied were of the clear cell type. Horse gram lectin, a marker of distal tubules, decorated this tumour type but only weakly. Concanavalin A and wheat germ lectin, which stained many renal tissue components, including proximal and distal tubules, also reacted with renal carcinomas.

The differential lectin staining of mesenchyme in various renal tumours may be important. Horse gram lectin and concanavalin A bound to mesenchyme only in Wilms' tumours, whereas winged pea lectin decorated mesenchyme in BMRTC (Table 2). It has been suggested that the cell substratum is important in the regulation of cell proliferation and morphogenesis. In an earlier paper we showed major differences in the expression of fibronectin and laminin among renal tumours. The addition of fibronectin to tissue culture medium reduced the migration of BMRTC cells into collagen but had no effect on Wilms' tumour cells in vitro.

The use of lectins did not help adequately in establishing the histogenesis of childhood renal tumours. There could be several reasons for this. Firstly, in the course of tumour progression the carbohydrate profiles of cells can change profoundly. Holthofer et al found that although renal carcinomas possessed several characteristics of proximal tubules, they did not stain by any of the lectins that specifically reacted with proximal tubules of normal kidney. They concluded: "these probes (lectins) are of limited value in histogenetic analysis of renal carcinomas." Secondly, fetal kidneys used in our present lectin binding studies were from 14-21 week old fetuses. It is conceivable that the use of kidneys from younger fetuses may prove valuable in exposing the origin of renal tumours: a Wilms' antisera has been shown to react with Wilms' tumours and fetal kidneys only to 10 weeks' gestation. Similarly, the studies of Ekblom et al showed that fibronectin is present in primitive uninduced nephrogenic mesenchyme but that after its induction by ureteric bud it disappears. An equally plausible explanation is based on the work of Sarioila, who has studied the histogenesis of the kidney vasculature. He transplanted a vascular embryonic quail kidney on to chick chorioallantoic membrane and found that fully developed transplanted kidney's blood vessels possessed markers of chick and not of quail, an obvious conclusion being that during normal nephrogenesis blood vessels of developing kidney originate from surrounding tissues rather than nephrogenic mesenchyme itself. Thus could it be that some primary renal tumours may have extranephric origin?

The technical assistance of Joan Ashworth is most gratefully acknowledged.

References

Renal tissues and lectins


Requests for reprints to: Dr S Kumar, Laboratory, Pediatric Oncology, Christie Hospital and Holt Radium Institute, Wilmslow Road, Withington, Manchester M20 9BX, England.
Study of childhood renal tumours using peroxidase conjugated lectins.
S Kumar, T Carr, H B Marsden and P H Morris-Jones

doi: 10.1136/jcp.39.7.736

Updated information and services can be found at:
http://jcp.bmj.com/content/39/7/736

**Email alerting service**

*These include:*

Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

**Notes**

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/