Controversies and advances in the management of Wilms’ tumour

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Wilms’ tumour is one of the success stories of paediatric oncology with long term survival approaching 90% in localised disease and over 70% for metastatic disease. Although appearing relatively simple compared to other cancer treatment regimens, successful treatment of Wilms tumour requires meticulous attention to correct staging of the tumour and good communication between the paediatric surgeon, pathologist and oncologist. The controversy of whether pre-operative chemotherapy results in a reduced overall burden of treatment compared to immediate nephrectomy has been addressed by the recently closed UKW3 randomised trial. Challenges remain in identification of histological and molecular risk factors for stratification of treatment intensity to allow safe reduction in therapy and avoidance of late sequelae for the majority while leading to increased biological insights and ultimately novel therapies for the minority of high risk tumours. Genetic predisposition to Wilms tumour is conferred by several genes, some of which cause malformation rather than cancer and may be of low penetrance. The proportion of children with heritable disease is uncertain and there remains a need to collect data on the need for screening in this susceptible population.

Wilms’ tumour is eminently curable, even when it has metastasised. The commonest site of such metastasis is the lung, followed by lymph nodes and liver. Wilms' tumour rarely metastasises to bone, bone marrow, or brain. Treatment consists of chemotherapy with one to three different drugs (usually vincristine, actinomycin D, with or without Adriamycin) together with surgical excision of the affected kidney. Radiotherapy is also used where there is residual or spilt tumour in the abdomen or metastases.

The aim of most recent national and international clinical trials has been to reduce the overall burden of treatment and potential for late sequelae, while maintaining therapeutic efficacy. Large studies involving thousands of patients have been run by the National Wilms' Tumour Study Group (NWTSG) in North America and the International Society of Paediatric Oncology (SIOP), mainly involving European countries. There is a philosophical divide across the Atlantic in that the NWTSG believes it is vital to identify accurate tumour staging by immediate surgery, whereas the SIOP approach is to reduce the risks of immediate nephrectomy on these often massive tumours by using preoperative chemotherapy to shrink the tumour. Postoperative treatment is then stratified according to the tumour staging information obtained at the time of delayed surgery. Both approaches result in similar tumour control rates but a different overall burden of treatment. In NWTSG studies, approximately 30% of patients have stage III tumours and therefore receive treatment with both an anthracycline, with its potential for permanent cardiotoxicity, and abdominal radiotherapy, with its adverse effects on growth and fertility. Using the SIOP preoperative chemotherapy approach, tumours are “down staged” so that only 15% of patients have stage III disease and 50% have stage I. However, SIOP currently treats patients with stage II disease with an anthracycline but no radiotherapy. Therefore overall, approximately 50% of children in SIOP studies receive an anthracycline but only 17% receive radiotherapy, compared to approximately 35% receiving both therapies in NWTSG studies. The proportion of children presenting with metastatic stage IV disease is similar in both studies. However, again, SIOP uses a response adapted risk stratification so that the two thirds of children with stage IV disease whose lung metastases resolve during the preoperative chemotherapy phase are not given pulmonary radiotherapy.

In an effort to further reduce the morbidity of treatment for Wilms’ tumour, the new SIOP WT 2001 study, which opened in the UK in March 2002, will ask a randomised question about the role of anthracyclines in intermediate risk stage II and III tumours.

THE UKW3 CLINICAL TRIAL

Over the past 10 years, the UK Children’s Cancer Study Group (UKCCSG) has run a randomised clinical trial comparing these two approaches. In this UKW3 study, all children with localised Wilms’ tumour deemed to be “operable” by the surgeon were eligible for randomisation to either immediate nephrectomy or biopsy and delayed nephrectomy following six weeks of preoperative chemotherapy. The only absolute criterion to render a localised tumour “inoperable” was documented tumour extension into the inferior vena cava.
cava. However, massive tumour size obscuring the renal hilum was a relative contraindication. The aim of this study was to compare the stage distribution using the two approaches with an anticipated 13% shift from stage III to stage I, without any impact on disease free survival. However, the study has not recruited at the anticipated rate because of a variety of factors. Many parents, when faced with the information that their child's tumour could potentially be removed on that day, preferred the surgeon to go ahead and remove it. On the other hand, many surgeons, when faced with a massive tumour that they know is very likely to shrink with preoperative chemotherapy, have elected for the latter approach. As a result, only one third of eligible children were randomised in the current study, with a majority of the non-randomised group being treated with immediate nephrectomy. The study closed to patient recruitment in March 2001. Providing that the shift in stage distribution from stage III to stage I is of the order of 15%, then even with reduced recruitment, the study should have sufficient power to make a significant contribution to the data. The first analysis is expected in late 2002.

ROLE OF BIOPSY
This is a somewhat controversial area as in the SIOP studies, if a tumour has the typical imaging and clinical features of Wilms' tumour, chemotherapy is started without histological proof. This results in approximately 1% of children with a non-cancerous renal lesion receiving chemotherapy. As actinomycin D carries with it an approximately 3% risk of hepatotoxicity, this was deemed unacceptable when the UKW3 trial was set up. Therefore, immediate biopsy was recommended for all children treated with preoperative chemotherapy. The outcome of this approach has recently been analysed and shows that 12% of renal tumours with the typical features of Wilms' tumour on imaging studies proved to be non-Wilms' tumour on prechemotherapy biopsy.4 As in the SIOP studies, 1% of children had a non-malignant lesion. There has also been controversy about whether a percutaneous cutting needle biopsy affects tumour staging and concerns about whether this could lead to an increased risk of flank recurrence. Reassuringly, the UKW3 approach has shown no evidence to suggest that performing such a biopsy should affect tumour staging or subsequent treatment.9

DURATION OF CHEMOTHERAPY
Both recent NWTS and SIOP studies have asked randomised questions regarding duration of therapy. The NWTS 4 study has shown that six months treatment is as effective as 15 months treatment in stages II to IV Wilms' tumour.1 The SIOP 9 study showed that four weeks was as effective as eight weeks of preoperative chemotherapy, and preliminary data from the recently closed SIOP 9301 study suggests that as little as four weeks postoperative chemotherapy is sufficient treatment for stage I tumours.1

TREATMENT OF STAGE I WILMS' TUMOUR
Selected stage I Wilms' tumours can probably be cured by surgery alone. The clinical challenge lies in identifying these patients at diagnosis. The currently running NWTS 5 study commenced by selecting patients less than 2 years of age at diagnosis with small tumours (nephrectomy specimen weight less than 550 g) and with a stringently defined tumour stage I (lacking any evidence of vascular or renal capsular invasion) for treatment with nephrectomy only. This arm of the study was closed in 1998 as the conservatively set early stopping rule for relapse rates was crossed. However, with longer term follow up, it seems that such an approach does not jeopardise overall survival as relapsing patients are highly salvageable. The UK has pioneered the treatment of stage I Wilms' tumour with minimal chemotherapy in the last three studies.1 These have shown that stage I Wilms' tumour (staged slightly more loosely than in NWTS 5) has a greater than 90% cure rate with only 10 weeks of vincristine monotherapy. As vincristine has very low acute morbidity and no documented potential for permanent late sequelae in children, this approach is very attractive and likely to be more widely adopted outside the UK. Currently, the NWTS treats stage I Wilms' tumour with two chemotherapy agents (vincristine and actinomycin D) for 18 weeks, whereas the SIOP approach uses the same drugs either side of nephrectomy for a total of eight weeks.

RISK STRATIFICATION
The ongoing challenge in all clinical trials is to identify better prognostic markers for stratification of therapy, reserving more intensive first time treatment for children with an anticipated high risk of treatment failure. The currently used criteria are tumour stage and pathology (see table 1). The NWTS 5 study is investigating the prognostic value of molecular markers, in particular allelic loss at a variety of chromosomal loci. It is predicted that tumours showing allelic loss at chromosome 1p, 16q, and/or 22q will be an adverse factor, but this requires confirmation in the current study. The SIOP therapeutic approach has the advantage that tumour response to preoperative chemotherapy can be examined for its utility as a prognostic factor. It appears, in addition to the well established presence of anaplasia as an adverse factor, that persistence of large amounts of viable blastema may also be adverse, whereas large amounts of necrosis and/or differentiation of the tumour into epithelial structures may confer a more favourable outlook. These hypotheses will be tested in the new SIOP WT 2001 study which the UK is joining.

GENETICS
Genetic predisposition to Wilms' tumour has long been recognised in the WAGR syndrome (Wilms' tumour, aniridia, genitourinary abnormalities, and mental retardation) and in certain overgrowth conditions, particularly Beckwith–Wiedemann syndrome (BWS) and hemihypertrophy.2 The genetic bases of these syndromes are becoming elucidated and have proven more complex than originally anticipated. The WAGR syndrome is caused by complete deletion of one copy each of the Wilms' tumour gene, WT1 and the adjacent aniridia gene, PAX6 on chromosome 11p13.13 Of interest, germ line point mutations in the same WT1 gene underlie the Denys Drash syndrome, a combination of early onset nephrotic syndrome, Wilms' tumour, and ambiguous genitalia.5 Both syndromes carry a high risk of Wilms' tumour of the order of 30%. More recently, similar germline WT1 mutations have been found in children with isolated nephrotic syndrome, particularly where renal biopsy shows focal glomerulosclerosis, and in Frasier syndrome, where the nephropathy has its onset in later childhood and there is predisposition to gonadal rather than Wilms' tumour.14 Most probably, these are all manifestations of a spectrum of abnormalities caused by constitutional WT1 mutation.13 The combined prevalence of Wilms'
tumour among all documented carriers of WT1 mutation suggests that the tumour penetrance is lower than previous estimates. The genetics of the overgrowth syndromes are more complex and they also carry a lower risk of Wilms’ tumour, no greater than 10%. BWS is a complex genetic disorder arising from mutations or abnormalities of imprinting in two or more genes in the 11p15.5 region. There is evidence to suggest that analysis of the methylation status of several genes in this region might predict an individual’s Wilms’ tumour risk. At a clinical level, it appears that those with documented nephromegaly in the first year of life are most at risk. However, before it can be advised that subgroups might be identifiable in which the risk of Wilms’ tumours is extremely low, this requires prospective evaluation. A registry of BWS patients and their tumours is maintained by Dr DeBaun in the USA and may provide information on this point in the future. Currently no such registry exists in the UK. In recent years it has become clear that other overgrowth syndromes may be associated with Wilms’ tumour. The Simpson Golabi–Behmel syndrome, an X linked overgrowth disorder with phenotypic overlap with BWS, has been shown to involve the GPC 3 gene. However, as yet no mutations in this gene have been found in sporadic Wilms’ tumour.

FAMILIAL WILMS’ TUMOUR

Familial Wilms’ tumour is rare but well documented, occurring in 1–2% of all cases of Wilms’ tumour. Usually such pedigrees are small, with only two or three affected relatives, and there is usually no associated congenital abnormality or predisposition to other tumour types. Genetic linkage studies in two of the uncommon large pedigrees has localised one gene for familial Wilms’ tumour, FWGT, to chromosome 17q. Another locus has been suggested at 19q, and a recent evaluation of all available pedigrees in the UK has confirmed that there is genetic heterogeneity for FWGT genes with at least four families clearly unlinked to any currently identified Wilms’ tumour locus. Until these genes are identified, it is difficult to predict what, if any, would be their involvement in sporadic Wilms’ tumour, both at the constitutional and somatic levels. However, the penetrance of FWGT at least appears to be low, of the order of 15–30%. It is therefore possible that a substantial proportion of apparently sporadic cases of Wilms’ tumour carry a constitutional mutation in a low penetrance familial Wilms’ tumour gene.

For the one Wilms’ tumour gene that has been isolated, WT1, there are clear clinicogenetic correlates. In a study of 201 cases of Wilms’ tumours selected from the NWTSG cases, only eight constitutional WT1 mutations were found, and these were virtually confined to boys with cryptorchidism as well as Wilms’ tumour (7/28 such patients). A separate study from Germany showed that somatic WT1 mutation was common in Wilms’ tumours showing stromal predominant histology (13/26) and remarkably, nearly all of these (10/13) had a constitutional WT1 mutation.

THE WT1 GENE

The WT1 gene was identified as lying within the constitutional deletions occurring in the WAGR syndrome. Although its structure suggests it functions as a transcription factor, identification of its physiological target genes has proved difficult. It is a complicated protein with alternative splicing that is clearly critical for normal development, as intronic mutations that prevent formation of certain splice isoforms are linked to genitourinary malformation and increased tumour risk. Following close on the heels of the retinoblastoma gene, it was initially anticipated that the WT1 gene would account for both heritable and sporadic forms of Wilms’ tumour. However, this turned out not to be the case, with no more than 5% of Wilms’ tumours being a result of constitutional WT1 mutation and a further 10% of sporadic tumours harbouring somatic mutations.

NEPHROGENIC RESTS

Nephrogenic rests are presumed precursor lesions for Wilms’ tumour. Their frequency of 1% in neonatal postmortem examinations compared with an overall instance of 1 in 10 000 children for Wilms’ tumour suggests that only 1% undergo malignant conversion. It is not yet clear how many genetic events are necessary for evolution from normal undifferentiated metanephrine blastema through nephrogenic rest to Wilms’ tumour. The role of chemotherapy in influencing this progression also remains unclear. However, it is currently suggested that prolonged treatment of up to a year’s duration with vincristine and actinomycin D may be of benefit. Nephrogenic rests can be subdivided into two types, intralobar and periblolar, which are thought to represent mutations occurring at an early or late stage of nephrogenesis respectively. Of interest, the former occur mainly in association with WT1 mutated Wilms’ tumours, whereas the latter occur mainly in association with BWS and hemihypertrophy.

SCREENING

Some children with specific malformations can be identified as being at high risk of Wilms’ tumour, of the order of 30%. With the discovery of the underlying genetic defects, in some cases subgroups can be defined which require screening. For example, in sporadic aniridia, high resolution karyotyping using probes for the contiguous PAX6, calmodulin, and WT1 genes on 11p13 can distinguish those children whose aniridia is a result of mutation confined to PAX6 and therefore not requiring screening from those with a more extensive deletion involving the WT1 gene. Similarly, children with early onset nephrotic syndrome involving diffuse mesangial sclerosis, even without ambiguous genitalia, are likely to harbour a constitutional WT1 mutation and hence carry an increased risk of Wilms’ tumour. In aniridia and BWS, where the risk of Wilms’ tumour is of the order of 10–30%, it is a generally held view that some sort of screening programme is justified. It has not been possible to perform randomised studies in this setting. However, retrospective analyses of the tumour stage at
LONG TERM FOLLOW UP

Children who survive more than three years from diagnosis of their Wilms' tumour are unlikely to suffer a recurrence, and the vast majority are at very low risk of developing second cancers related to their treatment. The aim of long term follow up is therefore to monitor renal function. The current UKCCSG recommendations for any child having a nephrectomy is that blood pressure should be checked annually and serum creatinine measured five yearly. An early morning urine should be tested annually for protein:creatinine ratio.

It is of interest that recent data from the NWTSG showed that children with Wilms' tumour and aniridia and also those with intralobar nephrogenic rests have a high incidence of renal failure with long term follow up beyond 20 years (fig 1). It is therefore important that such information is imparted to their adult physicians or general practitioners.

Analysis of the molecular genetics of Wilms' tumour has provided and continues to provide a fascinating insight into the relation between developmental abnormalities and embryonal cancers. Identification of genes involved in such processes and their impact on tumour biology may ultimately allow us to select safely subgroups of children with Wilms' tumour requiring only minimal therapy. The avoidance of anthracyclines and radiotherapy for an increasing majority would be a major step forward in the successful treatment of this tumour type. The new SIOP WT 2001 trial, which opens in the UK during 2002, will address this issue.

APPENDIX: USEFUL WEBSITES

Beckwith Wiedemann support network: http://www.geocities.com/bwss/index.html
UK Children's Cancer Study Group (UKCCSG): http://www.ukccsg.org
WT1 Mutational Database: http://www.umd.nekcer.fr

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