LETTERS TO THE EDITOR

Unilateral multicystic kidney

EDITOR,—Webb et al justify their policy of performing nephrectomy in healthy infants with prenatally detected multicystic dysplastic kidney (MCDK) on the basis of their findings in three children and their interpretation of the literature.1 Paediatricians may wish to scrutinise the methodology and conclusions of this paper more closely before referring asymptomatic infants for ‘prophylactic’ nephrectomy. On the evidence presented, the diagnosis of hypertension in their first two patients has not been conclusively proved. Furthermore, it is difficult to see how the renal pathology in their third patient can be attributed to MCDK in the absence of cysts and, more importantly, ureteric atresia.

Making a diagnosis of hypertension in children is problematic—indeed obtaining reliable reproducible blood pressure readings in fractious infants is very difficult. The measurement of peripheral plasma renin activity, which was performed in these children, is a relatively poor guide to the aetiology of hypertension. A credible diagnosis requires selective renal vein renin sampling.

The role of echocardiography in the diagnosis and assessment of mild or moderate hypertension in childhood has yet to be validated. The authors have not identified the source of the normal ranges quoted—an important consideration since different studies have yielded different ‘normal’ values for the parameters quoted and the ‘normal’ range has often been derived from a limited number of individuals in different age groups.

In their third patient, a hypertensive girl of 14, retrograde pyelography is said to have revealed ‘remnant renal tissue in the right loin’. Atresia of the pelviureteric junction or a blind ending ureter is a characteristic feature of MCDK which, on retrograde pyelography, is associated with the distinctive finding of a blind ending ureter. In this case the flow of contrast across the patient ureter into a remnant of the collecting system, effectively excludes the diagnosis of MCDK.

It is disappointing that the authors felt it necessary to quote the outdated findings of an informal postal survey of American paediatric urologists2 to support their arguments for nephrectomy. Published in 1978 and spanning an undefined period, this survey predated the era of prenatal ultrasound and many of the criteria demanded of a scientific study. In addition to misquoting the number of respondents to the survey (136 not 48), the authors have omitted to inform their readers that 10% of the paediatric urologists surveyed reported serious operative complications resulting from nephrectomy for MCDK including deaths, renal failure, intestinal obstruction, and removal of the wrong kidney.

The experience of other centres in relation to the risk of hypertension in MCDKs is different to that reported by Webb et al. Data on 441 children with MCDK was submitted to the Multicystic Kidney Registry from 49 centres in the United States and Canada. Of the 441, 260 were managed conservatively with follow up of up to five years in some cases. No cases of hypertension were reported in this series.

Ricklewood et al reported a series of 44 children with prenatally detected MCDKs.3 Blood pressure was measured continuously over a mean follow up of three years in 38 of these children and none was found to be hypertensive. These findings mirror our experience in Leeds where we have not encountered any case of hypertension associated with prenatally detected MCDK nor has an MCDK been identified in a child presenting with hypertension.4

Analysing 454 cases of hypertension admitted to the renal unit at the Hospital for Sick Children in Great Ormond Street, from 1975–85, Deal et al documented a wide spectrum of underlying renal pathology without encountering a single MCDK (J E Deal, personal communication).

A retrospective study of 42 children undergoing nephrectomy for renal hypertension in Glasgow did not include a single case of MCDK.5 Similarly, a detailed review of 22 cases of surgically treated hypertension of renal origin in childhood, undertaken in Boston, did not document a single MCDK.6

It is important to distinguish clearly between other forms of renal dysplasia and MCDK; a radial distinctive anomaly characterised by, non-communicating cysts and absent or non-functioning renal parenchyma, which is almost invariably associated with complete ureteric atresia or severe distal ureteric obstruction. The authors’ description of the radiological and pathological features of their third case clearly indicate that the renal pathology belongs in one of the former categories of dysplasia rather than representing a genuine MCDK.

Cases of hypertension conclusively linked to MCDKs have very rarely been reported but published evidence indicates that the magnitude of risk is extremely low. Menster et al, reviewing the published literature on the risks associated with prenatally detected MCDKs, concluded ‘It no longer seems advisable to routinely remove MCDK in young patients with either diagnostic or prophylactic reasons’.7

Paediatricians working in centres in which conservative management is undertaken can be reassured. The weight of published evidence continues to favour a non-operative approach to the management of prenatally detected MCDKs in healthy infants.

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Dr Webb and colleagues comment: Mr Thomas and Dr FitzPatrick raise some important questions that we are grateful for the opportunity to respond to them in detail.

We agree that the diagnosis of hypertension may be difficult in small infants and children, though having detected systolic blood pressures in excess of the 95th centile for both age and height on repeated occasions using an appropriately sized cuff with the patient in the resting state, we have met widely accepted criteria for the diagnosis of hypertension in all three cases.7 The presence of end organ damage (left ventricular hypertrophy diagnosed using accepted normal ranges8) in the first two patients confirms the longstanding duration and severity of this hypertension. It is generally agreed that raised peripheral plasma renin activity, while never truly diagnostic, is a useful indicator of renally mediated hypertension; renal vein sampling may be potentially technically difficult from the small vein which drains a MCDK and is generally used to predict response to surgery rather than to diagnosis it.

The resolution of hypertension and echocardiographic changes after nephrectomy provide further supportive evidence for the hypertension in these three children being renally mediated.

MCDK represents the extreme of a spectrum of obstructive cystic dysplasia; while strict pathological criteria require the presence of an atretic ureter for a diagnosis of MCDK to be made, in practice, physicians and surgeons make a clinical diagnosis on the basis of DMSA and ultrasound findings. Many children with lesions such as those seen in our third case may therefore be underfollowed with a clinical diagnosis of MCDK.

Our interpretation of page 213 of Bloom and Brosman’s paper9 was that 48 (35%) of the 136 urologists sent questionnaires responded to the section on clinical findings associated with MCDK, that 6 (15%) of 136 is a much larger number of urologists reporting hypertension than 15% of 48, and lends further weight to our argument supporting hypertension as a significant complication of the MCDK. Operative morbidity and mortality have fallen significantly since Bloom and Brosman’s publication; our previously reported absence of perioperative complications10 has continued over recent years.

The report of the Multicystic Kidney Registry on 441 children with MCDK in the United States and Canada of whom 181 underwent nephrectomy predominantly between 7 and 12 months of age,1 confirming that nephrectomy continues to be a valid treatment option in many centres.

Contrary to Thomas and FitzPatrick’s statement, four children followed up conservatively developed hypertension, though this was ‘believed to be unrelated to the multicystic kidney’. Without an alternative explanation or details explaining why hypertension was unrelated, MCDK related hypertension cannot be excluded. Our paper reports three further cases of hypertension in children diagnosed as having MCDK in addition to those others previously reported in the literature.

The surgical versus conservative management of the MCDK will remain an area of great controversy. Only by collecting complete and accurate data from large, unselected series of patients treated both surgically and conservatively will the true incidence of complications of both approaches emerge, and we make a further plea for the setting up of a national registry to expedite this.


EDITOR—Webb et al continue the debate about conservative versus surgical management of multicystic dysplastic kidney (MCDK) with case reports of three patients for whom surgery was undoubtedly indicated. While we would, however, caution against the experience of tertiary referral centres being extrapolated to the general population.

Experience at Northwick Park Hospital, a large district general hospital, suggests that MCDK is more common than the published incidence figures of one in 4300 live births.1 We have reviewed all cases of MCDK diagnosed at Northwick Park Hospital in the 10 year period 1987–96 during which there were 33 537 live births. During that period, routine antenatal ultrasonography was carried out at 16–18 weeks’ gestation. Further scans were performed at a later stage in the pregnancy only if there was a clinical indication. Of the 14 cases of MCDK diagnosed on antenatal ultrasonography, 11 were detectable on the routine scans. The gestation, including one case in which the pregnancy was terminated at 18 weeks because of contralateral renal agenesis and multiple congenital abnormalities. In those with a normal early scan, the abnormality was detected by ultrasonography at 25, 26, and 39 weeks’ gestation, respectively. One further MCDK was detected by palpation of a renal mass on the right side in a boy aged 6 months. The abnormality was detected by ultrasonography at 25, 26, and 39 weeks’ gestation, respectively. One further MCDK was detected by palpation of a renal mass on the right side in a boy aged 6 months. The abnormality was detected by ultrasonography at 25, 26, and 39 weeks’ gestation, respectively. One further MCDK was detected by palpation of a renal mass on the right side in a boy aged 6 months.

Gastrostomy in children with Crohn’s disease

EDITOR—We read with interest the article by Cosgrove and Jenkins, describing their experience with percutaneous endoscopic gastrostomy (PEG) in children with Crohn’s disease.1 In the past, we reported improved growth and overall safety of gastrostomy procedure in 16 children with Crohn’s disease.2 We have since had additional experience with seven children with Crohn’s disease and gastrostomy. The mean age of children was 13.3 years (range 2.5–17 years). Five had PEG and two had surgical gastrostomy tubes placed. The indication for gastrostomy was growth failure and refusal or failure to use nasogastric tube feeding. Follow up after gastrostomy placement was 1–28 months (mean 11 months). Elemental formula (Vivonex, Sandoz Nutrition) was used to provide 50–80% of required energy as continuous nocturnal infusion.

All children showed weight gain and an increased energy level. An increased growth velocity was noted in two children, in one from <1 cm/year to gastrostomy to 6 cm/year and in the other from 3.5 cm/year to 5 cm/year. Three children, two previously on nasogastric tube feeding, maintained their linear growth. 2 of the children were transferred from enteral nutrition (one and two months) was too short to assess the effect on linear growth in two children. Minor local complications including discomfort, granulation tissue, and local skin infection were observed in four of the seven children. Accidental removal of gastrostomy tubes occurred in two adolescent children. Closure of the gastrostomy tract after removal was complete in four children; three children still have the tube. The procedure and hyperalimentation were tolerated well, even by a 2.5 year old child.

Combining data from two studies1 and these seven children, most of the complications of gastrostomy were minor and easily treatable in a total of 33 children with Crohn’s disease. Two complications were noteworthy. Cosgrove and Jenkins reported one patient who developed intestinal obstruction by a leftover gastrostomy flange, a complication which is avoidable when early removal is done by endoscopy. In one child from our series, surgical closure was required for a persistent failure of gastrostomy tract to close. Gastrocutaneous fistula is a recognised complication after the gastrostomy procedure and was reported in 3.6% of patients without Crohn’s disease.3 More studies are required to assess whether the presence of gastric Crohn’s disease increases the risk of this complication.

Out of the 33 children with Crohn’s disease and gastrostomy, 26 (78%) showed improvement in linear growth and clinical wellbeing was obvious in majority. Although the number of children is still small, it appears that the advantages of gastrostomy outweigh the risks. However, we recommend that nasogastric tube feeding should be the first choice for supplemental nutrition as 25% of our patients with Crohn’s disease and growth failure used this method successfully for long time (mean duration 10.5 months).4 When this method is not possible, gastrostomy is a good and safe option. In case of previous abdominal surgery, surgical gastrostomy tube should be considered rather than PEG.

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Early micturating cystourethromgrams after urinary tract infection

EDITOR—Craig et al showed that timing of micturating cystourethrogram (MCU) does not affect the detection and grading of vesicoureteric reflux (VUR) if the test is performed more than one week after diagnosis and treatment of a urinary tract infection (UTI).1 There is a view that early MCU may reveal a higher prevalence of VUR due to transient inflammatory changes at the level of the vesicoureteric junction.2 This is to some extent refuted by the work of Craig et al,1 however their study was unable to address the issue of VUR in the first week after diagnosis of UTI when the possibility of such a situation would seem greatest.

Further to the issue of transient VUR, it is assumed that its presence is insignificant, therefore justifying a policy of delaying MCU (whether for one week or six weeks). The logic which supports this strategy may be questioned. An opposing view could be that if
UTI increases the likelihood of VUR at the time when the urine is infected, this may put the kidneys at risk in the same manner as more permanent VUR. Furthermore, such VUR will go undetected and the patient may not receive chemoprophylaxis. If such a child has another UTI, she or he may develop further temporary VUR putting the kidneys at risk once again. Perhaps this mechanism explains some of the renal scarring found in children investigated after UTI with normal MCU findings.

The issue of transient VUR needs to be progressed by further studies such as that of Craig et al, looking at MUC in the first week after UTI diagnosis, but not necessarily with the aim of dismissing transient VUR if it is found.

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Use of sucrose as a treatment for infantile colic

EDITOR.—In his commentary Dr Bell refers to ‘gripe water’ as being out of fashion. On the contrary, high street pharmacists will attest to the common use of gripe water for colic and other infantile ailments. In a survey of 200 infants in Shefield 64% received gripe water, 28% for 15 days or more. In the UK the sucrose content of gripe water varies between 10 and 20%. The dramatic response to 12% sucrose that was noted by Dr Markestad will illustrate many of the renal scarring. In: Holliday MA, Barratt TM, Avner ED, eds. Pediatric nephrology. Ed. Baltimore: Williams and Wilkins, 1994: 368.

Yeld 64% received gripe water, whereas many, such as those of scarlet fever, are of little value.


The eagerly awaited 15th edition of Recent Advances in Paediatrics has just arrived. There is no disappointment. Professor David has maintained the high standard we have been led to expect. This is an excellent series which enables paediatricians to keep themselves up to date with medical progress in a relatively painless fashion. This book is well worth buying and reading carefully.

The format is unchanged. There are a dozen chapters of general paediatric interest, and the comprehensive literature review by the editor. Each topic is clearly presented with helpful tables and numerous subheadings. Each finishes with a number of key points and numerous references. The subjects covered are wider ranging including bronchiolitis, large airways disease, thalassaemia, congenital hypothyroidism, and coeliac disease. There is a stimulating and partly speculative account of superantigen problems. Two chapters are devoted to problems in the newborn, pulmonary hypertension, and necrotising enterocolitis. A chapter on cholera, not usually a clinical problem in the UK, illustrates many of the health and social problems of the underdeveloped world. Spread may be rapid, the number of patients enormous, and the mortality appalling with 10,000 deaths in South America alone over three years. Good hygiene and clean water could prevent much of this, but governments lack the will to deal with poverty and the environment while population expansion and wars compound the problem.

Recent Advances in Paediatrics usually includes a chapter on recent progress in a physiological topic and its clinical application, and this is about nitric oxide. This is certainly a new topic which has only attracted study in the last few years, although apparently active biologically for ‘up to half a billion years’.

A chapter on headaches guides us through the appropriate imaging, stressing how uncommon it is to find unexpected pathology with a normal clinical examination, but recognising the role played in generating investigations. The fact that headaches and a brain tumour are extremely rare, are nearly always associated with other signs and symptoms, and get progressively worse should help paediatricians to resist reaching for the imaging request forms, with careful follow up being more appropriate.

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Imagine you will receive many similar letters—plus ça change...


BOOK REVIEWS


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The final chapter is on sleep problems. Before reading this my view was rather limited to children who don’t sleep and others who sleep too much, with occasional night terrors or seizures, and wondering when I will see my first case of narcolepsy. I’ve always been haunted by Professor Illingworth’s view which blames refusal to go to sleep on parental mismanagement and advises that the child be ‘left to cry’. I prefer William Cam’s advice that ‘The devil gets into them [babies] at an early age and of course you should have the baby in your bed’. At usual the best advice is probably somewhere in between the two extremes. This is an interesting chapter and should be read.

Finally, the excellent literature review. The main interest of course is ‘am I quoted?’ For me with only an occasional publication, this is unlikely, but in this case I am, slightly obliquely. Can you spot it?

MARTIN MONCRIEFF
Consultant paediatrician


Modest both in size and price this book stands between the grand reference book on paediatric infectious disease ‘Feigin and Cherry’ and the pocket sized manuals now available. Its challenge is to be a good, advanced reference book and yet be brief enough to be of use to the practising paediatrician who needs to look up a subject quickly. In many ways it meets the challenge. Much of the text is organised by systems rather than by infectious agents. Thus we are treated to chapters on respiratory, cardiac, gastrointestinal, and neurological infections, and those in the eyes and skeleton. Following these are sections on fever, neonatal infections, and HIV.

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because neonatal infection is included in the text. Investigation of such infection is written without reference to important viral infections occurring in the neonate. The index—one of the most important parts of any textbook—is usually neglected by the editors because they are exhausted or exasperated. Not all reactions to foods are immunological. Many are due to dietary elimination of foods that have been thus proved to trigger an adverse reaction. Lacking, however, are controlled trial data to evaluate this strategy.  

Almost every chapter ends with a suggested diagnostic or therapeutic trial for the reader, although there is no method of evaluation of these trials. Literature is not critically reviewed, and the chapters on asthma in children and adults respectively have much in common. There is almost no cross referencing between chapters. Without editing, the book could be considerably shorter.

Good points include liberal use of boxes containing key points, a generally readable style, and a healthy scepticism towards unproved and unscientific tests and treatments. There are helpful explanations of underlying theories of allergic disease, good descriptions of diagnostic tests including skin testing, and a comprehensive description of childhood asthma, which goes beyond what one might expect in a book confined to allergic disease.

There are thorough, although rather repetitive, chapters on allergic disorders of the upper respiratory tract, eyes, and skin. Weak points include paucity of illustrations, poor quality black and white photographs and, in the chapters related to drug treatment, there are too many pharmaceutical companies and not enough on trials of clinical efficacy. Surprisingly, in spite of the comprehensive account of asthma, there is very little on inhaler devices and nothing on comparisons of many foods containing toxic treatments.

However, the main factor limiting the usefulness of this book to British doctors is its North American parochialism. Thus the majority of the book is thus a series of essays on topics related to allergy and asthma. It makes an enjoyable read for anyone interested in this complex childhood endocrine disease but it is not a practical management manual. Only two of the chapters deal with any practical aspects of care, those on diabetic ketoacidosis and neonatal diabetes. The chapter on diabetic ketoacidosis discusses the physiology of the metabolic derangement in depth and includes the recent recommendations for its management from the British Society for Paediatric Endocrinology and Diabetes. The chapter on neonatal diabetes includes some useful pointers on management of this rare condition and an insight into its relationship to glucose intolerance in later life.

The majority of the book is thus a series of 'state of the art' dissertations from renowned authors with an emphasis on science and potential clinical applications rather than current practice. There is an excellent series of three review chapters on epidemiology, genetics, and possible preventive interventions. The paediatric aspects of the Diabetes Control and Complications Trial are crystallised in a multiauthor chapter headed by William Tamborlane, which also poses some interesting questions on how to achieve optimal management. The issue of pathogenesis and screening for microvascular complications is taken up in a chapter by Henrik Mortensen with, as might be expected, the emphasis on microalbuminuria.

There are also excellent chapters on maturity onset diabetes of the young, associated immunological diseases, and a final chapter on growth hormone and insulin-like growth factor-I and their role in the difficulties of adolescent management and possible therapeutic role in the future by Zvi Laron. This book should be essential and stimulating reading for all those paediatricians
involved in the care of children with diabetes. Not that it will greatly improve our care of them in its own right but it will go a long way to explaining much of the frustration in management and put all the important science behind the clinical facade. It provides a start to the ongoing care of these children and a hope that with just a little more knowl-edge more effective management, or possibly prevention, may not be too far away.

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Reviewing a textbook as detailed as this is indeed a daunting task. My initial response to the request was a look of disapproval ‘Why?’ but on reflection, having studied the book, I decided that a paediatrician in a district general hospital was exactly the right person to review such a tome. The layout and approach is novel and as such challenges the reader. Assessment based on a symptom or sign is a very practical approach. This provides a sense of security that all diagnoses have been considered when dealing with an unusual problem. However, that the cross referencing was not as detailed as I expected. For example, in an infant presenting with abdominal pain and vomiting, intussuscep-tion is only mentioned under vomiting, although pain is more common. Similarly, constipation warrants a detailed section of its own but it is not mentioned as a possible cause of chronic abdominal pain. Despite these difficulties, I enjoyed browsing through the book and I could not think of a symptom or sign that was not covered in depth. I particularly enjoyed the orthopaedic reviews of gait disorders and back pain which were both factual and user friendly.

My main criticism is that the print font is too small. Some of the tables and flow diagrams are very complicated, hence visually disconcerting. I would undoubtedly have opened and closed the book with one glance had I come across it in a bookshop. This would have denied me access to a wealth of information which in the severely disabled child can be overwhelming. Consultants may find it useful to refresh the memory of detail can be overwhelming. Consultants may find it useful to refresh the memory of detail can be overwhelming. Consultants may find it useful to refresh the memory of detail can be overwhelming. Consultants may find it useful to refresh the memory of detail can be overwhelming. Consultants may find it useful to refresh the memory of detail can be overwhelming. Consultants may find it useful to refresh the memory of detail can be overwhelming. Consultants may find it useful to refresh the memory of detail can be overwhelming. 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and the effects of reflux. The interlocking nature of problems is demonstrated by chapters on the consequences of malnutrition on neurodevelopment and the effects on the respiratory system of neurological deficit. Then the authors move to the assessment of a child with disability and feeding difficulties, a process focusing on strengths as well as weaknesses, clinically, nutritionally and by investigation. Two problems which greatly worry parents are refluxing and constipation, receive a chapter each. Therapeutic options in the disabled child are considered over two chapters and then the authors round off with personal views on the ethics and implications of feeding programmes for the child with the disability.

If the reader wishes to know how many nerves are involved in swallowing or about feeding teams, about Sandler’s ‘head cocking’ syndrome or the management of reflux, then this volume will help. The authors have made their ‘object’ improving the quality of life of the individual and the carer. This book will, with humanity and erudition, assist other in that aim.

A D KINDLEY
Consultant paediatrician


This is No 30 of Oxford Monographs on Medical Genetics. It discusses various demographic features of Arabs, selected disease entity among Arabs, genetic disorders in Arab countries and geographic regions, as well as cultural and religious attitudes to genetic issues. It provides an explanation of the important observation that there are many reports of genetic diseases in Arab populations. This could be explained by the large families and high level of consanguinity which increase the frequency of autosomal recessive conditions. In addition, with control of infectious diseases, genetic disorders will become a more prevalent problem.

Until now, the exact size of the problem of genetic diseases was not known because of lack of epidemiological population-based studies. However, there are several case reports and hospital based studies that indicate an increasing number of genetic diseases. Some of these were described first in Arab populations, for example, limb/pelvis, hypoplasia/aplasia syndrome described in a Palestinian family from Kuwait; subsequently reports came from other countries including Brazil and Italy.

No source of information has collected these reports in a systematic way, hence the importance this book which helps researchers in getting information on genetic diseases among Arab populations.

We can expect more description of genetic diseases in different Arab countries with advanced health care and improved laboratory technology. Therefore, revision and updating of this book are expected in the future because the subject is changing rapidly.

As a paediatric neurologist, I found the book very useful, in particular the chapter on new syndromes first reported among Arabs. More than 100 new single gene syndromes are described. Chapters on genetic disorders among the bedouin and cultural aspects are important for all doctors treating patients with genetic disorders. The chapters are based on the geography of Arab countries resulting in some repetition.

The editors have tried to cover the subject as broadly as possible but I think that coverage for certain diseases was not optimum, for example, endocrine and kidney disorders, congenital adrenal hyperplasia and congenital nephrosis, which are observed in certain Arab tribes. Similarly, the coverage of genetic and metabolic disorders in Saudi Arabia was also not complete, since Saudi Arabia is the origin of all the tribes of the other Arab countries. Many inborn errors of metabolism have been described from Saudi Arabia, for example in a special supplement of Developmental Medicine and Child Neurology in April 1991.

Despite these limitations the book provides a good source of information regarding genetic diseases in Arab countries. It is a good reference book for paediatricians, geneticists, endocrinologists, and other medical specialists in Arab countries as well as physicians from other countries who may face a medical problem in Arab patients and who may not be aware that these rare genetic diseases do occur in Arab populations frequently.

AHMED AL JARALLAH
Paediatric neurologist


There are few, if any, books related to paediatric endocrinology. The book is, I believe, a recommended unequivocally. Given the pace at which paediatric endocrinology has expanded, notably because of very rapid progress in molecular biology, textbooks involving this subject become quickly obsolete, therefore posing a greater challenge to the authors. This Colour Atlas of Paediatric Endocrinology and Growth tackles the subject courageously and with a different approach, mainly focusing on clinical aspects supported by lavish photographs. Unexpectedly, the text is thorough covering in detail the presenting signs, symptoms, differential diagnosis, and investigations while being very practical and avoiding superfluous information. It is written in a way which is easy to understand. Accompanying tables, clear growth charts, and appendices are simple to use and the most successful aspect of the book. They will certainly help unravel the subject and will act as a useful tool to memorise what ought to be retained by the reader. The illustrations act as the main lure for the buyers and have tried to them effective visualisation of diseases and concepts described.

The paragraphs regarding treatment end pathophysiology are deliberately brief but clear. The book’s intended audience is junior doctors as well as the general paediatrician who can easily extract practical information during daily practice. It is unfortunate, and yet understandable with the quality and the quantity of the illustrations that this small book is expensive. It would have been helpful if the authors had provided a few key scientific references at the end of each chapter. To sum up, in spite of minor misgivings, this book is a publication of great distinction; practical, ambitious, but simple and without a hint of self importance.

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Warning: do not take this title literally. Not only is this obviously not a paediatrician, but it is unlikely to fit in your pocket. However take the title figuratively, and it aptly describes this manual of paediatrics: complete but succinct.

Another handbook. Is there any need for it? And if so, is it any good? The answer to the first question depends on what the book aims to do and whether it has been done before. Written by the paediatricians of the British Columbia Children’s Hospital in Vancouver this book is based on the handbook given to their trainees and residents, and it aims to give practical advice to junior doctors, medical students, and nursing staff. The key word here is ‘practical’: a basic knowledge of the common conditions encountered in paediatrics is assumed, allowing it to focus on diagnosis and specific management. To the best of my knowledge this handbook is unique in this respect, and it is exactly what junior doctors require. Other handbooks either are restricted to major emergencies or a short version of a textbook, with non-specific or no advice on management.

However, for the same reasons I would question the usefulness of this book for medical students, at least in this country. The average medical student would be more likely to want to know what the haemolytic uraemic syndrome is, than how to manage it. That apart, the answer to the first question of whether there is any need for this book, is yes.

To answer the second question I put the book to the ultimate test and took it on call with me for a whole weekend. It was very useful. The chapters are clear and concise, with emergency management separated from background information and further management. Common conditions are given priority, and they are approached as they would be clinically, without having a final diagnosis. For example the cardiology chapter is divided into broad headings of cardiac failure, cyanosis and murmurs, rather than into individual structural cardiac conditions.

There is a comprehensive pharmacology section which includes common drugs and their major side effects and interactions of each drug, as well as a good chapter on surgical problems. The answer to the second question of whether the book is good, is also yes.

One final area which sets this book apart from its competitors is that it attempts to tackle some of the ethical and practical problems of being and surviving as a junior doctor. The authors obviously considered
these areas as important as how to manage a severe case of croup—important enough to include them in a pocket book. This is an attitude which is more prevalent on the other side of the Atlantic than it is here, and it is refreshing to see the issue being dealt with head-on, rather than relying solely on hard learned experience.

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TRAINING PACKAGE

Equal Rights/Equal Access—Improving the Care of Minority Ethnic Children with Disability or Chronic Illness. A Training Package. Produced by the Department of Child Health, University of Wales College of Medicine, Cardiff, 1996. ISBN 1-899717-05-6. The package is available from Joanne Plummer, Department of Child Health, University of Wales College of Medicine, Heath Park, Cardiff CF4 4XN, price £105.75. Anyone wishing to buy the package must attend the ‘Equal Rights/Equal Access’ course; for further information contact Joanne Plummer at the above address or on phone 01222 743375.

As paediatricians consider making services more child and family friendly, we come up against political issues such as inequality of care, the needs of minorities, and different cultural views. Having trained and worked in South Africa, I am only too aware of the impact such issues have on health care provision. The authors’ viewpoint is that we live in a multicultural society and need to be aware of people’s cultural background, at the same time recognising that cultures are mixing and changing and that all individuals are different in what has shaped their lives. We have a responsibility to take account of these factors in our work.

This manual was produced under the auspices of the Access to Black and Minority Ethnic Children with Disability Project in South Glamorgan. It details a six hour training course designed to be run by two trainers who between them have experience in training, race awareness, and disability in childhood. The emphasis is on promoting the understanding of multicultural practice, the impact of racism on service delivery, and the development of sensitive and non-judgmental attitudes. It is designed for professionals from various disciplines working with children with disability.

The authors have creatively and sensitively put together a format on how to run the course as well as the subject matter, allowing some flexibility to take account of local circumstances. Such a package of planning details, handouts, and overheads would be most welcome in other teaching that I do. The shifts in the course from discussion to group activities and video clips with suitably ethnic music, kept my attention and interest.

I particularly liked some of the examples relating to my own experience including the doctor asking for the child’s ‘Christian’ name; the father who does not believe that a woman paediatrician can be the ‘real doctor’; exercises highlighting our prejudices or expectations when we think we are taking ethnic issues into account and the trap of cultural stereotyping. The handouts raise issues such as poverty, and particularly pleasing is that they offer solutions, such as a comprehensive strategy for health service personnel including recommendations for managers and commissioners.

Reviewing this manual from the material provided without having participated in the training, is a bit like deciding to buy a car based on the manufacturers’ sales pitch, without having had the chance to test drive the car. These manufacturers may highlight the background research and the de-luxefeatures but the vital factor is to know whether the performance matches the specifications. I do however feel strongly tempted to buy.

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Archives of Disease in Childhood—http://www.archdischild.com

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