Our aim was to develop guidance for general paediatricians and primary care physicians in diagnosing and managing cow’s milk protein allergy in infants. The guidelines were developed by discussion based on existing national recommendations and standards, clinical experience and, whenever possible, evidence from the literature. Separate algorithms cover breast-fed and formula-fed infants. The recommendations emphasise the importance of comprehensive history taking and careful physical examination. Patients with severe symptoms need to be referred to a specialist. Elimination of cow’s milk protein from the infant’s or mother’s diet and challenges are the gold standard for diagnosis. This guidance is intended as a basis for local discussion, implementation and prospective evaluation. The algorithms should be regularly assessed using clinical audit standards. Once validated, the diagnostic framework could provide a standardised approach in epidemiological and therapeutic studies.

Between 5% and 15% of infants show symptoms suggesting adverse reactions to cow’s milk protein (CMP), while estimates of the prevalence of cow’s milk protein allergy (CMPA) vary from 2% to 7.5%. Differences in diagnostic criteria and study design contribute to the wide range of prevalence estimates and underline the importance of an accurate diagnosis, which will reduce the number of infants on inappropriate elimination diets. CMPA is easily missed in primary care settings and needs to be considered as a cause of infant distress and diverse clinical symptoms. Accurate diagnosis and management will reassure parents. CMPA can develop in exclusively and partially breast-fed infants, and when CMP is introduced into the feeding regimen. Early diagnosis and adequate treatment decrease the risk of impaired growth.

CMPA results from an immunological reaction to one or more milk proteins. This immunological basis distinguishes CMP allergy from other adverse reactions to CMP such as lactose intolerance. CMPA may be immunoglobulin E (IgE) or non-IgE associated. In IgE-associated cases, CMPA may be a manifestation of the atopic diathesis. In 170 unselected infants with a mean age of 7 months (range 2–11 months) with CMPA diagnosed by means of double-blind, placebo-controlled challenge, 58% showed an early reaction within 2 h after the last challenge dose. These early reactions usually manifest as urticaria, angio-oedema, vomiting or an acute flare of atopic dermatitis. The remaining 42% showed a later reaction, typically of atopic dermatitis or the gastrointestinal tract. Infants with early reaction were more likely to have a positive skin prick test (SPT; wheal size ≥3 mm) or test positive for specific IgE than those with later reactions. The amount of cow’s milk that elicited the immediate reactions varied from one drop to 161 ml.

In a selected group of 100 children with CMPA (mean age of 16 months), Hill et al reported that 27% developed symptoms, mainly urticaria and angio-oedema, within 45 min after ingesting cow’s milk. This represents the IgE-associated reaction. About half the children in this cohort showed pallor and gastrointestinal symptoms (vomiting and diarrhoea) between 45 min and 2 h after ingestion. The final 20% developed atopic dermatitis, respiratory symptoms or diarrhoea after more than 20 h and up to several days after the ingestion of cow’s milk. The proportion of children with early and late reactions, or positive or negative for specific IgE depends on how the patients were selected.

CMPA persists in only a minority of children. The prognosis (ie, the likelihood of becoming tolerant to CMP) depends on the patient’s age and titre of specific IgE at the time of diagnosis. In the experience of the taskforce members, children with proven CMPA who are radioallergosorbent test (RAST) or SPT negative become tolerant to CMP much earlier than atopic children with positive test results. Furthermore, patients with a history of IgE-positive CMPA are at increased risk of developing atopic diseases, such as asthma, atopic dermatitis and rhinoconjunctivitis, than those who were IgE-negative. Children with negative tests are less likely to develop multiple food allergy. Therefore, it is preferable to test for specific IgE (if not performed during the diagnostic work-up) in children with CMPA proven on challenge.

There are guidelines for the use of dietary products for the prevention and treatment of CMPA. However, there are currently no guidelines that specifically assist primary care physicians and general paediatricians in the accurate diagnosis and management of CMPA. This document aims to...
EVALUATION OF AN INFANT WITH SUSPECTED CMPA
A comprehensive history (including a family history of atopic) and careful physical examination form the foundation of both algorithms. The risk of atopy increases if a parent or sibling has atopic disease (20–40% and 25–35%, respectively), and is higher still if both parents are atopic (40–60%). In comparison to cow’s milk formula-fed infants, exclusive breast feeding during the first 4–6 months of life reduces the risk for CMPA and most severe allergic manifestations during early infancy. 17 The distinction between breast-fed (fig 1) and formula-fed infants (fig 2) reflects the importance of ensuring an adequate duration of breast feeding. Management principles also differ. The management of breast-fed infants depends on reducing the maternal allergen load and strict avoidance of CMP in supplementary feeding. It is recommended that exclusive or partial breast feeding is continued, unless alarm symptoms (table 1) require a different management. The earlier CMPA develops, the greater the risk of growth retardation. 18

Unfortunately, there is not one symptom that is pathognomonic for CMPA. The most frequent symptoms of CMPA are listed in table 2. The timing and pattern of these symptoms aid the differential diagnosis. Symptoms of CMPA occur often, but not always, within the first weeks after the introduction of CMP. Many children with CMPA develop symptoms in at least two of the following organ systems: gastrointestinal (50–60%), skin (50–60%) and respiratory tract (20–30%). 1 The symptoms associated with CMPA can range from mild to moderate to severe, although this stratification is by its nature subjective. In this guidance, symptoms that put the child at an immediate risk of growth retardation (such as failure to thrive or growth faltering) differentiate severe from mild-to-moderate CMPA. 15

Differential diagnoses include, among others: metabolic disorders, anatomical abnormalities, coeliac disease and other (rare) enteropathies, pancreatic insufficiency (such as in cystic fibrosis), non-immunological adverse reactions to food (such as fructose malabsorption or secondary lactose intolerance, mostly with an onset in older children), allergic reactions to other food allergens (such as hen’s eggs, soy, wheat, etc) or other substances (such as animal dander, moulds, dust), malignancy, and infections (particularly gastrointestinal and urinary tract infections) and sepsis. A role for allergy in recurrent otitis media has been heavily discussed in some of the literature. 20

The clinician should also assess whether the child suffers from concurrent conditions. For example, 15–21% of children with suggested or proven gastro-oesophageal reflux disease (GORD) or CMPA suffer from both conditions. Furthermore, 16–42% of children with a history of GORD show signs or symptoms of CMPA. CMPA has also been related to infantile colic. However, colic has numerous aetiologies which should be considered during the differential diagnosis. However, there is a subgroup of about 10% of colicky formula-fed infants in whom the colic episodes are a manifestation of CMPA. 21

While in some young infants there is a strong association between atopic dermatitis and CMPA, many cases of atopic dermatitis are not related. The strength of the association depends on the age and severity of the atopic dermatitis: the younger the infant and/or the more severe the atopic dermatitis, the stronger the association. 22

Reactions to other foods, especially egg and soy, but also wheat, fish, peanut and other foods depending on the regional dietary intake, may occur in combination with CMPA. Therefore, complementary feeding and, preferentially, all supplementary feeding should be avoided during the diagnostic elimination diet.

ALGORITHM FOR THE DIAGNOSIS AND MANAGEMENT OF CMPA IN EXCLUSIVELY BREAST-FED INFANTS
Breast feeding is the gold standard for milk feeding in infant nutrition and is recommended exclusively for the first 4 months of life at least. The incidence of CMPA is lower in exclusively breast-fed infants compared to formula-fed or mixed-fed infants. Indeed, only about 0.5% of exclusively breast-fed infants show reproducible clinical reactions to CMP and most of these are mild to moderate. This might be related to the fact that the level of CMP present in breast milk is 100 000 times lower than that in cow’s milk. 23 In addition, immunomodulators present in breast milk and differences in the gut flora in breast-fed and formula-fed infants may contribute to the prevalence of CMPA in breast-fed compared to formula-fed infants. The most frequent symptoms of CMPA in exclusively breast-fed babies are listed in table 2 and include general dermatological and gastrointestinal manifestations. Severe forms of CMPA (table 1) are very rare in exclusively breast-fed infants. The occasional cases that occur are usually severe atopic dermatitis with protein losses and failure to thrive. Other rare conditions suggesting severe CMPA include anaemia due to colitis with rectal bleeding and protein-losing enteropathy. In these cases, introducing CMP into the infant’s diet (eg, supplementary feeding) may exacerbate the symptoms. Cases with alarm symptoms should be referred to a paediatric specialist for further diagnostic work-up and management. In these infants, diagnoses other than CMPA are much more likely, and identifying the correct diagnosis should not be delayed.

Breast feeding should be promoted for the primary prevention of allergy, but breast-fed infants with proven CMPA should be treated by allergen avoidance. There is evidence that food proteins from milk, egg, peanut and wheat are excreted in breast milk and may cause adverse reactions during exclusive breast feeding in sensitised infants. Due to the many benefits of breast feeding to the infant and the mother, clinicians should advise mothers to continue breast feeding but avoid the causal foods in their own diet. Egg avoidance studies indicate the foetus may be exposed to maternally-derived egg antigens despite maternal dietary avoidance measures. 24 In infants with atopic dermatitis, the risk of being sensitised to milk was four times higher, and to egg eight times higher, than in infants without atopic dermatitis. 25 Age at first introduction of solid food and diversity of solid food showed no effect on atopic dermatitis incidence. 26 However, there are no data on additional systematic elimination of hen’s egg in symptomatic infants.

Therefore, as fig 1 shows, if the infant develops symptoms of allergy, a maternal exclusion diet avoiding food containing...
 CMP and hen’s eggs is advised by the task force although the evidence for CMP is more exhaustive than for hen’s egg. In a subgroup of children with severe atopic dermatitis, peanut could as well be eliminated from the mother’s diet since peanut allergy is more likely in children with atopic dermatitis. When deciding which foods with a high allergenic potential to suggest avoiding (hen’s eggs rather than, for example, wheat and fish), the taskforce considered evidence that in most geographical regions egg proteins are the most common cause of allergy after CMPA in infants and young children. The evidence that peanut allergy can cause severe symptoms has been well established, but not in exclusively breast-fed infants. In contrast to milk and egg, peanut consumption is common in only parts of the world such as the USA, UK and some other European countries. In primary prevention, which is not the topic of this manuscript, it has been shown that peanut is secreted into breast milk following maternal ingestion.29 Since peanuts are not an essential nutritional part of a normal diversified diet, they are easy to avoid, and since infant sensitisation through breast feeding has been suggested, the task force suggests eliminating peanut as well from the mother’s diet (although the evidence for peanut is much weaker than for cow’s milk and egg). The task force recognised the difficulties in implementing such widespread dietary recommendations. Further studies are required to test the feasibility of such programmes and whether they are effective if implemented on a large scale.

Furthermore, a diet that also excludes fish, wheat and other gluten-containing grain products is very demanding for the mother and may increase the mother’s risk of consuming an unbalanced diet. Therefore, the relative risk associated with an extensive, first-line exclusion diet may be greater than the potential benefit. In a secondary approach, the additional elimination of wheat and fish will require the advice of an experienced dietician in order to ensure that an adequate nutritional intake is maintained. If the mother has a certain suspicion that another food elicits the symptoms in her child, the elimination diet should be adapted accordingly. In some very rare cases, such as in infants with severe atopic dermatitis with impaired growth, breast feeding should be stopped.18 However, the authors strongly propose that these infants should be referred to a specialist before breast feeding is discontinued.

The elimination diet should be continued for a minimum of at least 2 weeks, and up to 4 weeks in cases of atopic dermatitis or allergic colitis. The mother will require calcium supplements (1000 mg per day divided into several doses) during the elimination diet. If the elimination diet fails to improve the symptoms, the mother should resume her normal diet and a referral to a specialist should be considered, depending on the type and severity of the infant’s symptoms.

If symptoms improve substantially or disappear during the elimination diet, one food per week can be reintroduced to the mother’s diet. If symptoms do not re-appear on reintroduction of a particular food to the mother’s diet, the elimination of that specific food can be discontinued.

If symptoms re-appear, the food responsible should be eliminated from the mother’s diet as long as she is breast fed.

Figure 1 Algorithm for the diagnosis and management of cow’s milk protein allergy (CMPA) in exclusively breast-fed infants. eHF, extensively hydrolysed formula.
feeding. If solid foods are introduced into the infant’s diet, care should be taken to ensure solids are free from the food proteins that the infant is allergic to. If CMP is the responsible allergen, the mother should continue to receive calcium supplementation during the elimination diet. If the mother is on a CMP-elimination diet for a long period, appropriate nutritional counselling is required. When the mother wants to wean her infant, the child should receive an extensively hydrolysed formula (eHF) with demonstrated clinical efficacy.

ALGORITHM FOR THE DIAGNOSIS AND MANAGEMENT OF CMPA IN FORMULA-FED INFANTS

Patients with life-threatening, particularly respiratory symptoms or anaphylaxis, conditions need to be referred immediately to an emergency department experienced in the treatment of this condition. In all the other situations, the initial step in the diagnostic work-up for CMPA is clinical assessment accompanied by history taking, including establishing whether there is a family history of atopic disease (fig 2).

The algorithm differs according to the severity of symptoms (fig 2). If the infant does not present alarm symptoms (as listed in table 1), the case is considered as mild-to-moderate suspected CMPA, and a diagnostic elimination diet should be initiated. Infants presenting with symptoms such as angio-oedema of lips and/or eyes, urticaria and immediate vomiting are likely to have IgE-mediated allergy. In the case of IgE-mediated allergy, improvement (and normalisation) offers a safety net before challenge. A positive SPT increases the likelihood of a positive food challenge but not the severity of the reaction. In the study from Celik-Bilgili and coworkers, 60% of the patients with a RAST class 1, 50% in class 2, 30% in class 3 and even 20% in class 4 had a negative food challenge.30

DIAGNOSTIC WORK-UP IN SYMPTOMATIC INFANTS WITH NO ALARM SYMPTOMS (MILD-TO-MODERATE MANIFESTATIONS)

In a case of suspected mild-to-moderate CMPA, CMP elimination should start with a therapeutic formula for CMPA. The guidelines define a therapeutic formula as one that is tolerated by at least 90% (with 95% confidence) of CMPA infants.31 These criteria are met by some eHFs based on whey, casein or another protein source, and by amino acid-based formulae (AAF). Preferentially, all supplementary food should be stopped during the diagnostic elimination diet. If this is not possible in infants beyond 6 months, only a few supplementary foods should be allowed with dietary counselling. Nevertheless, the diet should not contain CMP or hen’s eggs, soy protein or peanut. Referral to a paediatric specialist and dietary counselling may be needed for patients who do not improve. In such cases, further elimination of other allergenic proteins such as fish and wheat may be appropriate. In most cases, the therapeutic elimination diet should be given for at least 2 weeks, although this may need to be increased to up to 4 weeks in gastrointestinal manifestations and atopic dermatitis before deciding that the intervention has failed. eHFs that meet the definition of a therapeutic formula are the first choice. An AAF is indicated: if the child refuses to

Figure 2  Algorithm for the diagnosis and management of cow’s milk protein allergy (CMPA) in formula-fed infants.
diagnosis should be considered. Higher cost. If symptoms do not disappear on the AAF, another specific gastrointestinal manifestations or both. In these cases, clinicians should refer to a specialist for further diagnostic work-up.

Children may react to residual allergens in eHF, which may be one reason for the failure. The residual allergens in eHFs seem to be more likely to produce gastrointestinal and other non-IgE-associated manifestations compared to AAFs. However, IgE-related reactions have also been reported with eHF. In such cases, clinicians should consider an AAF which has been proven to be safe and nutritionally adequate to promote weight gain and growth. In some situations, the infant may be initially switched to an AAF, especially if they experience multiple food allergies, specific gastrointestinal manifestations or both. In these instances, the potential benefits of an AAF may outweigh its higher cost. If symptoms do not disappear on the AAF, another diagnosis should be considered.

The role of in vitro and in vivo testing for CMPA

Table 1: Alarm symptoms and findings (can be found alone or in combination with items listed in table 2), indicating severe CMPA as the possible cause

<table>
<thead>
<tr>
<th>Organ involvement</th>
<th>Symptoms and findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gastrointestinal tract</td>
<td>Failure to thrive due to chronic diarrhoea and/or refusal to feed and/or vomiting</td>
</tr>
<tr>
<td></td>
<td>Iron deficiency anaemia due to occult or macroscopic blood loss</td>
</tr>
<tr>
<td></td>
<td>Hypoalbuminaemia</td>
</tr>
<tr>
<td></td>
<td>Endoscopic/histologically confirmed enteropathy or severe colitis</td>
</tr>
<tr>
<td>Skin</td>
<td>Exudative or severe atopic dermatitis with hypoaalbuminaemia or failure to thrive or iron deficiency anaemia</td>
</tr>
<tr>
<td>Respiratory tract (unrelated to infection)</td>
<td>Acute laryngoedema or bronchial asthma with difficulty breathing</td>
</tr>
<tr>
<td>General</td>
<td>Anaphylaxis</td>
</tr>
</tbody>
</table>

Table 2: Most frequent symptoms of CMPA

<table>
<thead>
<tr>
<th>Organ involvement</th>
<th>Symptoms</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gastrointestinal tract</td>
<td>Frequent regurgitation, vomiting, diarrhoea, constipation (with/without perianal rash), blood in stool, iron deficiency anaemia</td>
</tr>
<tr>
<td>Skin</td>
<td>Atopic dermatitis, swelling of lips or eye lids (angio-oedema), urticaria unrelated to acute infections, drug intake or other causes</td>
</tr>
<tr>
<td>Respiratory tract (unrelated to infection)</td>
<td>Runny nose (allitis media), respiratory infection, wheezing</td>
</tr>
<tr>
<td>General</td>
<td>Persistent distress or colic (wailing/irritable for &gt;3 h per day) at least 3 days/week over a period of &gt;3 weeks</td>
</tr>
</tbody>
</table>

*Infants with CMPA in general show one or more of the listed symptoms.

www.archdischild.com
During oral provocation the dose of formula should be titrated as follows. After a physical examination of the undressed infant, with inspection of the skin, a drop of the formula is put on the lips. If no reaction occurs after 15 min, the formula is given orally and the dose is increased stepwise (0.5, 1.0, 3.0, 10, 30, 50 to 100 ml) every 30 min. Thereafter, the infant is observed for 2 h and examined for cutaneous and respiratory reactions before going home. If no reaction occurs, the child should receive at least 250 ml of cow’s milk-based formula each day for the next week and the parents told to observe the child for late reactions.

Positive challenge: CMPA confirmed
If symptoms of CMPA re-appear, the suspected diagnosis of CMPA is confirmed and the infant should be maintained on an elimination diet using eHF or AAF until the child is between 9 and 12 months of age, but for at least 6 months, whichever occurs first. The challenge is then repeated. If it is possible to follow the infant with IgE-mediated allergy with SPTs and/or specific IgE determination, normalisation or improvement of these tests would help in choosing the time point of challenge. Supplementary feeding should be introduced carefully to avoid accidental intake of CMP. Nutritional counselling must ensure a sufficient intake of the therapeutic formula (eHF or AAF) to guarantee adequate calcium intake.

Negative challenge: no CMPA
Children who do not develop symptoms on the cow’s milk formula during challenge and up to 1 week after follow-up can resume their normal diet, although they should be monitored. Clinicians should advise parents to be attentive for delayed reactions, which may evolve over several days following the challenge.

DIAGNOSTIC WORK-UP IN INFANTS WITH SEVERE MANIFESTATIONS
Formula-fed infants suspected of suffering from severe CMPA should be referred to a paediatric specialist. In the meantime, an elimination diet should be started and the child should preferably receive an AAF. AAF is recommended because infants in this group fail to thrive, suffer from macronutrient deficiencies or have pain. In these cases, AAF minimises the risk of failure on an eHF and further weight loss. Many of these children may need further diagnostic work-up to rule out other diagnoses. However, the recommendation to use AAF as a first choice is based on clinical experience, not on evidence. This approach should be prospectively validated.

The decision concerning allergen challenge in cases with severe CMPA should always be made by a specialist and performed in a hospital setting. In cases with a history of a life-threatening reaction, a food challenge may be contraindicated.

DISCUSSION
These recommendations have been developed as guidance for general paediatricians and primary care physicians to assist with the diagnosis and management of CMPA in breast-fed and formula-fed infants. They emphasise the importance of breast feeding, which is the preferred method of feeding healthy infants. The recommendations also underscore the importance of a comprehensive history taking (including a family history of atopy) and a careful physical examination to exclude other causes, identify any concurrent conditions and classify the condition as mild-to-moderate or severe CMPA. The algorithms differ according to the method of feeding (breast-fed or formula-fed infants) and according to the severity of symptoms. Blood-stained stool in an infant is alarming for the mother, although recent evidence suggests this is a benign and self-limiting phenomenon, mostly occurring in exclusively breast-fed infants. CMA in these patients is less common than previously believed, and an association with viruses can be observed in some patients. CM challenge is thus essential in infants who become symptom-free during a CMP-free diet to reduce the number of false-positive diagnoses of CMPA. In cases with recurrence of symptoms after reintroduction of dairy products in the mother’s diet, the algorithm recommends eHF if the mother wants to start weaning the infant and if the child is younger than 9–12 months. However, one could speculate that since the infant reacted to the (very) small amounts of proteins present in its mother’s milk, it might be preferable to recommend AAF. Unfortunately, no data are available on this topic. Patients with severe symptoms need to be referred to a specialist experienced in managing childhood allergies.

In formula-fed infants, clinicians should consider whether SPTs, patch tests and determination of specific IgE would aid the diagnostic work-up and guide management. However, elimination diets and challenges are the gold standard for diagnosing CMPA in formula-fed infants. For simplicity and for socio-economic reasons, an open challenge is recommended by the taskforce. In the case of a doubtful outcome, a double-blind placebo-control challenge is helpful. If a reduction in the cost of diagnostic testing is important, RAST, SPT or both can be limited to those infants responding to an elimination diet to guide the challenge or after a positive challenge to predict the prognosis more accurately.

Infants with mild-to-moderate symptoms should receive eHFs, or AAF if the infant refuses to drink eHF or if the cost–benefit ratio favours AAF, for at least 2–4 weeks. Children who show a substantial improvement or disappearance of symptoms should undergo a challenge under medical supervision. If symptoms of CMPA emerge upon food challenge, the child should be maintained on eHF or AAF for at least 6 months or until 9–12 months of age. If symptoms do not improve on eHF, primary care physicians and general paediatricians should consider an elimination diet with AAF, other differential diagnoses or both for the symptoms and/or refer the patient to a paediatric specialist.

If the clinician suspects severe CMPA in a formula-fed infant, the patient should receive AAF and be referred to a paediatric specialist experienced in managing infant allergies. Food challenges in infants with severe symptoms should be performed only in a setting with personnel experienced in treating anaphylaxis. The clinician should be aware that severe reactions may also occur in patients with previously mild-to-moderate reactions after a period of dietary elimination.

The use of unmodified mammalian milk protein, including unmodified cow’s, sheep, buffalo, horse or goats’ milk, or unmodified soy or rice milk, is not recommended for infants. These milks are not adequately nutritious to provide the sole food source for infants. Furthermore, the risk of possible allergic cross-reactivity means that these milks or formulas based on other mammalian milk protein are not recommended for infants with suspected or proven CMPA.

Soy protein, for example, is not hypo-allergenic. The incidence of soy allergy in soy formula-fed infants is comparable to that of CMPA in cow’s milk formula-fed babies. Adverse reactions to soy have been reported in 10–35% of infants with CMPA, regardless of whether or not they were positive or negative for specific IgE antibodies for CMP. In particularly, infants with multiple food allergies and eosinophilic enterocolitis syndrome react to formulas which include soy protein.

Although soy formulations are significantly cheaper and have a better acceptance than eHF and AAF, the risk that the child will develop soy allergy in addition to CMPA, particularly in infants below 6 months of age, was considered by the authors to be too high for it to be recommended as the first choice. Soy may be considered in infants refusing to drink eHF and/or AAF, especially
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Funding: The consensus panel, the literature search and the drafting of the manuscript were funded by a grant from SHS/Nutricia. The paper was drafted by Mark Greener, a medical writer. SHS International Ltd and Nutricia did not have any editorial control over the final manuscript, which remains entirely the responsibility of the authors.

Competing interests: DH, CD, MB, SK and YV declare they have received support for clinical research projects from SHS/Nutricia and the same authors and MB declare they have presented lectures at SHS/Nutricia-sponsored meetings. Also, SK has presented lectures at sponsored meetings and received research support from Nestec S.A. and Nestle. YV has received support from Janssen Pharmaceuticals, Astra, Wyeth, Biocodex and Nestle. None of the other authors made any declarations relevant to the preparation of this manuscript. The authors declare the absence of competing interests and confirm their independence regarding the content of this manuscript.

Yvan Vandelenas and Sibylle Koletzko are joint lead authors.

REFERENCES

neurophysiology”. They emphasise that “neurophysiologic studies provide an important extension to the clinical evaluation and are predicated on a careful history and examination”, rather than being tests to interpret in isolation. They asked the contributors “to provide succinct descriptions of clinical disorders where neurophysiologic testing is a useful adjunct”. This pragmatic marriage of technical and clinical considerations shines through much of the text, and I feel that the editors have succeeded in their aims.

The 46 contributors are predominantly from North America, but there are four from Europe and two from Australasia. For the most part, the information is generic and, the chapter on the diagnosis of brain death, for example, has an orientation towards legal and practical issues pertaining to North America.

The book covers electroencephalography (EEG), evoked potentials (somatosensory, brainstem auditory and visual) and the clinical neurophysiology of the motor unit (electromyography and nerve conduction studies). It is divided into four sections: basic principles and maturational changes; disorders of cerebral function; neuromuscular disorders; and other neurophysiological techniques. This last section is relatively brief but covers magnetoencephalography (MEG), transcranial magnetic stimulation (TMS) and the assessment of sphincter dysfunction.

The first part of the book contains chapters describing the normal features of EEG in the neonatal and paediatric age groups. Separate chapters, divided into age periods, outline an approach to the visual analysis of EEG with clear and didactic suggestions about extracting essential features. This works well.

The second part of the book is devoted to the investigation of disorders of cerebral function. Having read chapters 2 and 3, which cover the features of the normal neonatal EEG and suggest an ordered approach to its visual analysis, one is faced with chapter 15, which describes the abnormal features of the neonatal EEG. This presents information in a logical, progressive and user-friendly manner, with clear reviews of normality and age-dependent changes that are separated from details of abnormal conditions and findings. It also includes information on abnormalities in the various forms of evoked potentials, and a chapter on their use in intra-operative monitoring.

There are excellent chapters on childhood sleep-wake disorders, drug effects, infectious diseases, trauma, and metabolic, toxic and degenerative diseases.

The chapter on EEG in the evaluation of children for epilepsy surgery is very brief at a mere seven pages. However, epilepsy surgery is an immense area and, on balance, I think that the editorial decision to substantially restrict this section is reasonable.

The third part of the book again takes a usefully clinically oriented approach, with chapters on the floppy infant, facial and bulbar weakness, disorders of the anterior horn cell, plexopathies and radiculopathies, and focal neuroptides. There are substantial chapters on autonomic testing in various conditions, including Guillain-Barré syndrome, chronic autonomic neuropathies, diabetes mellitus and neuromuscular transmission defects. A whole chapter is devoted to the relationship between DNA analysis and neurophysiological aspects of neuromuscular disorders. Given the relative frequency of exposure to neurophysiological examinations in these age groups, the book is weighted heavily towards discussion of neuromuscular disorders, but then this is a broad field with a large number of rare diseases that merit some coverage.

Unsurprisingly, it is possible to find minor points of imperfection that might be addressed in a second edition, for example inconsistent headings within a table on classification and a figure on scalp electrode positions that is poorly reproduced.

The book is well produced to a standard typical of Elsevier products. The text and pictures are generally black and white, although there are five colour plates. The index is large and comprehensive but is not divided into separate author and subject indices, and omits, for example, some important scoring systems mentioned in the text.

The book is competitively priced but, because of its specialist nature, it is unlikely to reside high on the wish list of any but the most enthusiastic general paediatrician. However, it is an excellent text for paediatric neurologists and neurophysiologists, particularly those in training. The orientation towards young people and the coverage, in one volume, of EEG, peripheral neurophysiological and other techniques makes it an efficient and very useful learning and reference text. It is an essential element of the clinical neurophysiology departmental library in any centre that performs these investigations in young people.

Andrew L Lux

Corrections

doi:10.1136/adc.2007.116608

E Fitzpatrick, B Bourke, B Drumm, et al. Outcome for children with cyclical vomiting syndrome (Arch Dis Child 2007;92:1001–4). In table 2 of this paper row “Medication prescribed”/column “Resolved” should read 8/25 (32%) (not 16/28/25 (32%) ). In addition, row “Trigger factor identified”/column “Resolved” should read 16/25 (64%) (not 25 (64%).)

doi:10.1136/adc.2006.110999corr1


In figure 2 of this article the arrow pointing to the right from the box “Open challenge; Cow’s milk formula under clinical observation” should actually point to the box “CMFA symptoms; Maintain CMF elimination diet until 9–12 months of age, and for at least 6 months” and not to the box “No CMFA symptoms; Resume CMF in diet and monitor” as published.

In addition, in figure 2 the box “Elimination diet” should have included the additional text: Therapeutic Extensive Hydrolysed Formula (eHF) for 2 to 4 weeks (*) .

doi:10.1136/adc.2007.115493


doi:10.1136/adc.2005.080721

S Friedman, S Reif, A Assia, et al. Clinical and laboratory characteristics of non e. coli urinary tract infections (Arch Dis Child 2006;91:845–6). The fourth author of this paper, Ram Mishaal, was inadvertently omitted. We apologise for this error.