FAY

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Dr E Sumner MA BM BCh FRCA

28th June 2004

Dear Dr. Sommer (for i formation

Thank you for your letter dated 11th June 2004, in which you express your great unease regarding the understanding of the basics of fluid management and their implementation in clinical practice in children and young people. As you copied your letter to Dr Campbell, CMO I felt it best to discuss the points you have raised with her before replying.

Indeed, in the interim, the results of a N. Ireland regional audit have also become available, assessing the implementation of the hyponatraemia guidelines issued by DHSSPS in early 2002. These show an encouraging level of compliance with the Xguidelines in paediatric units across the province, but do also identify some situations in which the guidelines do not appear to have been fully followed. It appears that this is not a problem unique to Northern Ireland, as shown by the attached letter published in Archives of Disease in Childhood in July 2003, relating to the death of a child from hyponatraemia in a major paediatric teaching hospital in England. We have also become aware of issues relating to the use of oral fluids and the potential for complications to arise when these are administered (often by parents) to children receiving IV fluids. These are often hypotonic as many children refuse to drink proprietary oral rehydration formulas, and we believe that this issue will also be worthy of further attention. In addition, concerns have recently been expressed by colleagues in adult specialties regarding care of children requiring intravenous fluids who come under their care, often in an adult environment.

In recognition of the concerns which have become apparent from all of these sources we feel that there are a number of actions which need to be taken. I understand that Dr Campbell will be making arrangements for a workshop at which issues of fluid management can be discussed between colleagues in relevant specialties within medicine, and indeed nursing. In addition, I have already highlighted with the General Medical Council the importance of specific reference to education and training in fluid administration and management for doctors in the PRHO grade, as part of the current

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revision of 'The New Doctor'. It will be helpful if the importance of this is also raised with the GMC by others, possibly including Mr Leckey and yourself. We will also bring this issue to the attention of the Northern Ireland Postgraduate Dean and Director of Undergraduate Medical Education, so that it can be raised with relevant individuals and committees who have responsibility for both undergraduate and postgraduate training.

When the audit results were presented in my own unit last week we agreed with our nursing colleagues that a formal morning and evening handover of fluid management involving relevant medical and nursing staff should be introduced for all children receiving intravenous fluids.

We are very grateful for the time you have given to helping identify these important issues, and guiding our thinking towards developing solutions. I hope that the steps set out above will show that this is a subject which the profession in Northern Ireland are taking very seriously, not just with the rapid development and circulation of the 2002 guidelines and the subsequent editorial in the November 2003 Ulster Medical Journal, but also with the regional audit which has subsequently been undertaken, and our plans to follow this up in the ways I have outlined. We will of course be delighted to hear of any other ways in which you feel we could usefully take this issue forward.

With best wishes,

Yours sincerely

Dr John Jenkins.

Senior Lecturer in Child Health & Consultant Paediatrician

cc Dr H Campbell CMO

Mr J Leckey HM Coroner

infiltrating angiolipama; a case presenting with Kasabach-Musritt syndrome. Arch Dis Child 2003;88:67-8.

2 Barlow CF, Priebe CJ, Muliken JB, et al. Spastic diplogio as a complication of interferon Alfa-2a treatment of hemangiamas of infancy. J Pediatr 1998;132:527-30.

3 Warls H. Maass E. Kohler B, et al. Interferon alpha-2a therapy in haemangiomas of inlancy: spastic diplogia as a severe complication. Eur J Padian 1999;158:344

4 Vesikari T, Nuutila A, Cantell K. Neurologic -sequelae following interferon therapy of juvenile laryngeal papilloma. Acta Pediatr Scand 1988:77:619-22

Ezekowitz RAB, Mulitken JB. Folkman J. Interferon alfa-2a therapy for life threatening hemangiomas of infancy. N Engl J Med 1992,326:1456-63.

6 Vial T, Descates J. Clinical taxicity of interferons. Drug Sal 1994:10:115-50.

### Authors' reply

Dr Biban states that we did not adequately emphasise the neurologic side offects of interferon treatment. Although it has been reported that interferon alpha has been responsible for various neurologic side effects, there fare no clear data indicating the frequency of drese in children. Sport term interferon therapy has been safely used at our departmont in treating various different conditions, particularly in the complex hemangiomas for many years. No side effects of interferon therapy except mild fever, malaise, leukope. nia, and elevation of liver transaminases have been observed. These were reversible by stopping therapy for a short period. In one patient who received long term interferon therapy, peripheral houropathy developed during the treatment

This patient was a 15 year old boy with Hodgkin's disease who received interferon as an adjuvant immunotherapy post autologous Stem cell transplant. Peripheral neuropathy developed 20 months after IFN treatment.' A large comulative dose combined with the prolonged treatment may have had an important role in this complication in our case. We concluded that the use of interferon in children differed by KSM or in didldren with various bénign tumours containing vascular elements is still a good therapeutic alternative. If the duration of treatment and the cumulative doses of interferon are closely monitorised, we recently cared for a 13 month old gir Therapy would not be an important problem. As the use of interferons in various conditions gradually expands, the data related to the adverse neurologic side effects will increase and be better understood.

5 Emir, C Akyüz / Deportment of Paedintics, Ankara University, Ankara, Turkey

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Emir S, Kulluk T. Chan KW ret al. Paripheral neuropathy during interferon therapy in a child with Hodgkin's disquise. Ped Hem Oncol! 1999;16:557

Acute renal failure and cystic fibrosis

It is surprisping that there are few imports of acute renal failure (ARF) in childred with cystic fibyosis (CF) given the large number of antibiotic courses prescribed and the passi-/ bility of either direct toxicity from aminoglycosides of the occurrence of interstitial ness

phritis. The registry of our regional paedlatri renal unit shows no cases of ARF in a C patient between 1985 and 1998, but thre cases between 1999 and 2001, all of whor had received gentamicin and ceftazidine.

Over the past nine months we have been referred three additional CF patients who has been treated with a combination of gen tamicin and ceftazidine/cefuroxime (toble 1) The initial doses of antibiotics used to trea the patient were within UK guidelines,' bu the gentamicin levels were raised. All six chil dren had received a number of other medica tions including, in some instances, other anti biotics prior to the gentamicin cephalosporin combination. Only one of the four biopsy specimens revealed interstitia nephritis in addition to the acute tubula. necrosis (ATN) changes found in all four Al six children have made a good renal recovery with normal blood pressures and creatining levels at three months.

A recent e-mail survey of members of the British Association for Paediatric Nephrology revealed four other cases of ARP with combination antibiotic therapy in CF patients (three of four with cestazidine and gentamicin). The increased incidence points to the need for increased vigilance when gentamicin and cephalosporin combinations are used to treat exacerbations, particularly if there is a potentially dehydrating state or pre-existing tenal anomaly. The cases have been reported to the Committee for the Safety of Medicines and we suggest a national monitoring programme should be instigated.

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> Correspondence lo: Dr A R Walson; Inoyes I @nohl.trent.nhs.uk

#### References

Drew JH, Watson AR, Evans JHC. et al. Amibialitz and acute renal fallure in children with cystic fibrosis. Poediatric Perinoial Drug Therapy 2002;5:65-7.

Cystic Fibrasis Trust. Antibiatic treatment for cystic fibrosis. Report of the Cystic Fibrosis Trust, Bromley, Kent, September 2002,

## Fatal iatrogenic hyponatraemia

admitted to hospital following a short history of diarrhoes and vomiting. Clinical examination revealed lethargy and moderate dehydration. Initial serum sodium was 137, mmol/l and she was commenced on intravenous fluids using 4% dextrose/0.18% saline.

Twelve hours after admission the child suffered a generalised tonic-clonic seizure at which time the serum sodium was found to be 120 mmol/s. Unfortunately, the child went on to have a respiratory arrest, developed fixed dilated pupils, and died despite full intensive care. An extensive postmortem examination revealed only diffuse cerebral swelling with necrosis of the cerebellar tonsils.

It is well recognised that symptomatic hyponatraemia can result in significant morbidley and mortality in previously healthy children' and adults.' The administration of hypotonic intravenous fluids to children can be fatal and the reasons for this have been well documented for several years. Many physiological stimuli encountered during actite illness result in the non-osmotic release of antidiuretic hormone, these include pyrexia, nauses, pain, reduced circulating vol. cine, and the postoperative state. The adminntration of hypotonic intravenous fluids in

PostScript

these circumstances results in the excretion of hypertonic urine, the retention of free water, and the development of hyponatraemia.

Despite clear and repeated warnings over the past few years. The mutine administration of 4% dextrose/0.18% saline remains standard practice in many paediatric units. This practice is based on formulas developed for calculating maintenance fluid and electrolytes in healthy children over 40 years ago and there seems little understanding of the potential risks associated with their use during acute illness.

A global change of clinical practice is required to prevent these needless deaths. This is a challenge that the RCPCH should face up to, together with the Medicines Control Agency and the National Patient Safety Agency. A useful first step would be to label bags of 4% dextrose/0.18% saline with the warning that severe hyponatraemia may be associated with its use.

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#### References

Arieff Al, Ayus JC, Froser CL. Myponokaemia and death or permanent brain damage in healthy children. BMJ 1992;304:1218-22.

2 Bhaila P, Edon FE, Coulter JBS, et al. Lesson of the week: hyponatroemic selzures and excessive intake of hypotonic fluids in young children. BWJ 1999; 319:1554-7

3 Sjablam E, Hajer J. Ludwigs U, et al. Falal hyponatraemic brain cedema due la common gastraenteritis with accidental water intexication. Intensive Care Med 1997;23:348-50.

4 Halberthal M, Halperin ML, Bohn D. Acute hyponatroemia in children admitted to hospital: retrospective analysis of factors contributing to its development and resolution. 8MJ 2001;322:780-2.

5 Bohn D. Children are another group at risk of hyponatraemia perioperatively. BAU 1999;319:1269.

O Durward A, Tibby SM, Murdoch IA.
Hyponotraemia can be caused by standard
fluid regimens. BMJ 2000;320:943.

7 Maritz Mt. Augusta

Moritz ML, Ayus JC, Lo Crosse encephalitis in children. N Engl J Med 2001;345:148-9.

# Thyroid screening in Down's syndrome: current patterns in the UK

Children and adults with Down's syndrome are at increased risk of developing thyroid dysfunction, and screening for thyroid dysfunction is recommended as part of their health surveillance.' Clinical history and examination are known to be unreliable indicators of myroid sysfunction in Down's syndrome Venous blood for thyroid stimulating humbone (TSH) assay remains the gold ndard. Capillary blood spot on filter paper

SH has been proposed as a simpler and more convenient alternative screening method for typothymidism in these children.

It is establish current screening practices, we under took a postal questionnaire of community paediatricians registered with the British Association for Community Child Health (BACCH). Community paediatricians are the group mostly likely to see children with Down's syndrome for health surveillance. Paediatricians were asked whether they routinely screened children with Down's syndrome for thyroid dysfunction. They were asked at what age of child they began screening, how often they screened and which method they used.

The questionnaire response fate was 64% (209/325). All the paediatrigians who returned completed questionnaires routinely looked after children with Down's syndrome. sexpected, almost all of respondents, 93% (194/209), were greening routinely. Most paediatricians began screening before 5 years of age, and screened every two years (table 1). Venous blood TSH was the most frequently used method of screening (83%, 174/209). Only a small number have begun using capillary blood spot on Mier paper TSH (7%, 15/209). A few paediamicians were relying on clinical suspicion alogic Those paediamicians not routinely screening for thyroid dysfunction, were either moasuring TSH opportunistically or were undertaking biochemical screening only when symptoms or signs raised suspicion.

The Down's Syndrome Medical Interest Group (DSMIG) has recommended biochemical screening for thyroid dysfunction at least every two years after the fifst year of life." Most paediamicians' practice is consistent with this recommendation dapillary blood sampling has practical advantages over venous sampling, with regard to patient acceptability, payticularly in adolescents with Down's syndrome and with regard to cust. There is growing evidence that capillary blood spot TSK is a reliable screening tool for thyroid dysfunction in children with Down's syndrome. Capillary blood spot Tou may, in the future, come to replace vendus TSH sampling in children with Down's syndrome.

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### References

DEMIG. Basic medical surveillance essentials for people with Dawn's syndrome.
Recommendations of the Dawn's Syndrome Medical Interest Group, Thyrold Disorder, DSMIG (UK) 2001, www.dsmig.org.uk.

2 Nobie SE, Leyland K, Findlay CA, et al. School based screening for hypothyroidism in Down's syndrome by dried blood spot TSH measurement. Arch Dis Child 2000; \$2:27-31.

3 Phillp M, Murphy J, Mayne P, et al.
5 Screening for hypothyroidism in children and adolescents with Down's syndrome. Arch Dis Child 2000:82 suppl 1):A16.

4 Varadkor S. Reynolds AP, Lessing D. Thyroid dysfunction in Down's syndrome: a comparative study of two screening methods.

Arch Dis Child 2000; \$2 [suppl 1]: A46.

# Changes in serum sodium levels during treatment of hyperglycaemia

Carlotti el al' state that fluid and electrolyte management might contribute to the development of cerebrai oedema in hyperglycae-l'imia. There is a simple rule of thumb, formulated by Karz, which may help calculate water and electrolyte deficits and predict the changes in sodium levels which accompany changes in glucose levels,' namely that a decrease of 0.29 mmol/l in serum sodium may be expected for every 1.0 mmol/l increment in serum glucose.

This may be explained as follows: hypergly-caemia causes an osmotic movement of water out of the cells, which leads to hyponatraemia by dilution. Thus, at presentation, the patient is usually severely dehydrated intracellularly. However, the serum sodium is lower than would be expected because of this dilution of the extracellular fluid. When the patient is treated with insulin, glucose enters the cells, taking water with it, leading to a relative concentration of the extracellular fluid, and thereby a rise in serum sodium. This rise may be predicted and calculated using Katz's formula.

Carlotti et al 2150 comment on the report of Glaser et al that the chance of cerebral oedema during treatment is increased in children who present with high initial serum urea levels and when there is a lack of an increase in serum sodium levels during treatment.' This increased risk may be explained by the fact that the uses level rises in proportion to the degree of dehydration. Urea contributes to serum osmolality and if the fall in urea is not taken into account the serum osmolality may be allowed to drop too rapidly, thereby increasing the risk of cerebral oedema. Car. lottl et al do not take this into account in their formula for calculation of osmulality. The calculation of serum osmolality as twice the sum of sodium and potassium plus the urea and glucose levels (all in mmol/l) corresponds better with the formally measured osmolality.\*

By treating hyperglycaemia using hypotonic solutions or glucose alone, the serum osmolality will fall rapidly and thereby increase the risk of cerebral oedema.

Serum asmolality must be monitored frequently, either by direct measurement or calculation from the sodium, potassium,

Age screening initiated (y)  c5 5-10 Voidate	No. (%)  167 (80%) 28 (13.5%) 1 (0.5%) 13 (6%)	Screening frequency  Yearly Two yearly Three yearly Opportunist celly Other No dale			٠
			No (%)	Screening method	No. (%)
			35 (17%) 115 (55%) 20 (10%) 17 (8%) 10 (4.5%) 12 (5.5%)	Venous TSH Capillary blood spot TSH Both venous and capillary blood spot TSM Clinical history and examination only No data	174 (83%) 15 (7%) 4 (2%) 3 (1.5%) 13 (6.5%)