Hyperpyrexia during Anaesthesia

SIR,—Malignant hyperpyrexia has now been classified into two groups: a rigid type and a non-rigid type.1 Malignant hyperpyrexia with rigidity should be easy to identify by the rigidity itself, serum muscle enzyme studies (during and after the event), and muscle biopsy. On the other hand, malignant hyperpyrexia without rigidity is virtually a diagnosis of exclusion, since this group of patients have normal serum muscle enzyme levels and a normal muscle biopsy.2 How then is non-rigid malignant hyperpyrexia to be distinguished from other pyrexial reactions during anaesthesia? The non-rigid hyperpyrexia-prone patients apparently fail to produce a neuromuscular block when given intravenous suxamethonium.3 This is the reason for the so-called testing test and requires voluntary co-operation by patients, which may not always be forthcoming. Since no simple test is available, a more critical approach should be made before classifying a pyrexial reaction as non-rigid malignant hyperpyrexia, or this group will be in danger of becoming a dustbin for all types of pyrexial reactions that occur during anaesthesia. Where there is an alternative explanation for a pyrexia occurring during anaesthesia a diagnosis of malignant hyperpyrexia should not be made.

Drs. Sheila Kenny and H. Rolfe (21 November, p. 492) report a case of hyperpyrexia during anaesthesia, which they classified as non-rigid hyperpyrexia when there might equally have been an alternative explanation. Their patient already had a preoperatively pyrexia, with presumably some peritonitis. The temperature may have started to rise between when it was last recorded and the beginning of the operation, especially if atropine or scopolamine pretreatment were given. The temperature would have risen further when heat loss was reduced by covering the patient in surgical drapes. The situation reminds one of the so-called “ether convulsions” which occurred in preoperatively pyrexial children with peritonitis. How easy was it to cool the patient—how much iced fluid was given? The fulminating rise of temperature in malignant hyperpyrexia is not usually easy to reverse. Since a diagnosis is being made by exclusion, the results of blood culture and/or peritoneal swab culture should be taken into account. Did the patient receive any antibiotic therapy after surgery? Both antibiotics and closing of the peritonaeum might have been sufficient to account for the return of temperature to normal on the next day. The metabolic acidosis could be a consequence of the rise in temperature (and Metabolic rate) during anaesthesia. While this may yet be a case of non-rigid malignant hyperpyrexia, it is not conclusively proved if an alternative explanation cannot be excluded.

All anaesthetists are now aware of the danger of malignant hyperpyrexia, and should treat any pyrexia occurring during anaesthesia expectantly. However, because they do this, it does not mean that all pyrexias are due to malignant hyperpyrexia.—I am, etc.,

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Rational Oxygen Therapy

SIR,—Dr. J. M. Leigh makes the acceptable plea that the dosage of oxygen during oxygen therapy should be better controlled (5 December, p. 620). May I add that in addition to using high air flow with oxygen enrichment, using the Ventimask series, the use on the wards of the hospital oxygen analysers provides a very practical method of monitoring the inspired oxygen during therapy with patients breathing spontaneously, and makes minute to minute adjustment easy. The method is flexible as it can be used in most circumstances when controlled oxygen dosage is of importance. If sampled via a fine nasopharyngeal catheter breath-by-breath concentrations can be measured. Oxygen analysers of the paramagnetic type are expensive and equipped to require the measurement of blood oxygen tension. They are suitable for use by nursing staff and the readings can be recorded with those of the usual parameters.—I am, etc.,

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Low Birth Weight

SIR,—Your leading article (19 September, p. 657) was of interest and in particular the reference3 to survival and subsequent development of infants weighing between 850 and 1,250 g. at birth. Results from this hospital on infants of similar birth weight distribution may be of interest.

During the four-year period 1966–69 65 infants weighing between 850 and 1,250 g. at birth were liveborn, of whom 32 (49%) survived. Subsequent follow-up—to one year in every case—showed evidence of permanent brain damage in 6 (18%) and a further three were considered suspect. During the same four years there were 19,844 livebirths, of whom 1,288 (6-5%) weighed 2,500 g. or less at birth. Neonatal death in the first week occurred in 134 (10-5%).—We are, etc.,

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Facial Paralysis after Local Dental Anaesthesia

SIR,—In their report on this very rare complication of local anaesthesia (24 March, p. 798) Leader I. B. Twining and Flight Lieutenant T. Keane (28 March, p. 798) list many causes of facial palsy. We report a further case, one of three seen this year in which facial paralysis followed local anaesthesia, because of one of the factors responsible for those that they mention may have been responsible.

A woman aged 25 years gave a life-long history of eczema, hay fever, and asthma. Three weeks before the relevant dental treatment she was unwell and enlarged painless glands appeared in both sides of her neck which persisted for five days. One hour after injection of local anaesthetic around a lower premolar tooth there was a rapid onset of partial bilateral ptosis, diplopia, and vertigo. The ptosis cleared within two days, but defective conjugate movement of the eyes in all directions with fine nystagmus on lateral gaze persisted. She lost her visual symptoms after four days, when a complete right lower motor neurone facial palsy, without aregia, and severe Rombergism supervened. These signs rapidly responded to treatment with prednisone and cortisone. Three weeks after the patient attended the A.N. and was cured. Her very relaxed facial expression and the fact that she included a number of punctures, was negative. Three weeks later she was completely cured.

Obscure neurological syndromes may be encountered in 1% of cases with infectious mononucleosis, and when the isolated cranial nerve is involved differential diagnosis may be difficult. In a recent report1 a patient with infectious mononucleosis appeared to be developing a posterior fossa tumour but it was the 5th and 6th cranial nerves that were involved and later the cerebellum progressively became involved, but this neurological condition was cured with the administration of steroids.

Our patient developed after dental anaesthesia lesions within the mid-brain and later the pons who was rapidly treated with steroid therapy. She had had some painless glanular enlargement in her neck earlier, and though we could not prove the diagnosis we suspected that this was infectious mononucleosis. However, in view of her long history of various forms of allergic response, flitting lesions in the brain stem of an angioneurotic oedematous type are a more likely explanation for her very dis...
pressing neurological disorder. In either case, it would seem that this patient’s brain stem activity had become set precariously and that the arrival of local anaesthetic caused its temporary breakdown.—We are, etc.,

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J. M. Norcliffe Roberts.

Reference


Carbenicillin and Hypokalaemia

Sir.—Brunner and Frick1 drew attention to the development of hypokalaemia and metabolic alkalosis with high dose sodium benzylenzepinylcillin therapy. Their patients received 60 g. daily for periods of 10 to 14 days. The standard dose of carbenicillin (α-carboxybenzylenzepinyl, Propen) for systemic pseudomonas infections is 30 g. per 24 hours. No serious electrolyte disturbances have been recorded to date in patients receiving this antibiotic.2 However, we have seen recently two patients who developed hypokalaemic alkalosis with carbenicillin therapy and wish to draw attention to this potentially dangerous complication.

An 80-year-old woman fell sustaining an undisplaced pertrochanteric fracture of her left femur which was treated conservatively. Urinary retention occurred, requiring an indwelling catheter. Pseudomonas pyocyanica were cultured in pure growth from repeated specimens of urine. She became febrile, had marked anorexia, vomited from time to time, and developed a painful, red, swollen wrist suggestive of bacteremia although blood cultures proved sterile. Investigations: serum sodium 139 mEq/l., bicarbonate 27 mEq/l., urea 29 mg./100 ml. Two days later Carbenicillin 30 g./24 hrs. was given in two litres of normal saline in view of her dehydration. Five days later she had sacral oedema and a raised jugular venous pressure. The infusion volume was halved and bendrofluazide 5 mg. per day given. The patient was afebrile a week later, and felt well, but was hypokalaemic: serum sodium 138, potassium 2.0, bicarbonate >30 mEq/l., urea 30 mg./100 ml. The carbenicillin was stopped and the hypokalaemia rapidly responded to oral potassium chloride.

A 12-year-old woman was involved in a road traffic accident sustaining a compound fracture of her left tibia and fibula with extensive soft tissue damage. An initial exploratory operation revealed injury to the posterior and posterior tibial arteries. The leg became infected and repeated swabs yielded pure cultures of Pseudomonas pyocyanica. The patient was febrile with spikes of fever up to 104°F (40° C.).

On 12 September carbenicillin was started, 5 g. 4-hourly into a dextrose saline infusion. The next day an above knee amputation was performed. Carbenicillin was stopped on 21 September when the patient was complaining of lethargy and weakness and was found to have sluggish tendon reflexes. A day later investigations showed a severe hypokalaemic alkalosis: serum sodium 131, potassium 1.5, bicarbonate 40+ mEq/l., urea 10 mg./100 ml. Again oral potassium chloride in large doses rapidly corrected the electrolyte disorder.

In the first case bendrofluazide and a poor dietary intake may have been partly responsible for the hypokalaemia. In the second case there were no such contributory factors. The mechanism of penicillin-induced hypokalaemia is not certain, but probably depends on penicillin acting as a non-reabsorbable anion thus increasing passive distal tubular potassium excretion down an increased negative transstubular potential difference.3 Although carbenicillin contains 4.7 mEq of sodium per gram, an increased distal tubular sodium load is probably not generally an important factor as sodium delivery is only rate limiting for the hypothetical cation exchange mechanism in certain circumstances.4 However, both the above patients had other factors tending to reduce sodium excretion with probable increased aldosterone production. The first patient developed heart failure, was found to have a serum albumin of only 2.2 g./100 ml. and was on phenylbutazone. The second patient had a general anaesthetic and surgery which are often followed by salt and water retention. Distal tubular sodium reabsorption may thus have been increased in both instances, encouraging tubular excretion of potassium especially in view of the relative deficiency of chloride.5

The manufacturers recommend that for severe infections probenecid be used to increase blood levels of carbenicillin. Probenecid was not used in the present cases and might well have prevented hypokalaemia by reducing renal tubular levels of carbenicillin. As there is no preparation of probenecid for parenteral administration it is likely that carbenicillin will be used on many occasions by itself. Carbenicillin is a valuable drug in the treatment of pseudomonas infections which were cured in both our patients. However, on occasions it can give rise to dangerously severe hypokalaemia.—We are, etc.,

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References

2 Price, J. D., personal communication.

Labelled Fibrinogen in Renal Transplantation

Sir.—Mr. J. R. Salaman (30 May, p. 517) has suggested that surface measurements may have particular value in recipients with delayed function of renal transplants in the first few weeks after transplantation, and that a percentage of transplant radioactivity of more than 120% of that of the heart would appear to indicate rejection. We have performed similar studies in nine patients here,1 and have found that late transplant rejection is accompanied by surface accumulation of fibrinogen radioactivity. However, it was observed that during the early phase after transplantation it was the wound healing rather than rejection of the transplant that accounted for the increased radioactivity. Thus, the patient shown in Fig. 1 had a marked increase of radioactivity over the transplant although there was no evidence of rejection, and there was a similar increase in radioactivity over the nephrectomy wound of the other side. Although in the patient shown in Fig. 2 the control side, where the nephrectomy had been performed two years before transplantation, showed no increase of radioactivity, the increase over the transplanted side was probably due to the process of wound healing, since there was no evidence of rejection. The bladder was emptied before each measurement and no haematomas were present. It may be noted (right half of both