concentration of up to 28 mmol/l has been shown not to interfere with measurement of serum amylase with this technique (personal communication, Boehringer Mannheim).

In view of his heavy alcohol intake, we considered that our patient may have chronic relapsing pancreatitis in which a normal serum amylase is seen more commonly. This is unlikely, however, as the patient was well between acute episodes, with no abdominal pain. In addition, there is no evidence of pancreatic exocrine insufficiency and no pancreatic calcification or other features of chronic pancreatitis on plain abdominal radiograph or abdominal CT scan. Finally, our patient has had no further episodes of pancreatitis since effective treatment of his hyperlipidaemia.

**Final diagnosis**

Recurrent acute pancreatitis with normal serum amylase concentration.

**Keywords:** pancreatitis, hyperlipidaemia


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**Recurrent dehydration in a young girl**

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A 50-day-old girl was admitted to hospital because of fever, vomiting and refusal of feeding for four days duration. She was born normally at term to healthy parents who were first cousins. On admission she was dehydrated and weak. Basic blood biochemical investigations were normal apart from a serum sodium of 159 mmol/l. She was given parenteral therapy with subsequent improvement of her condition and she was discharged in good health after a few days. She was readmitted again the following week with fever, lethargy and marked dehydration without an associated history of vomiting or diarrhoea. Investigations showed a serum sodium of 165 mmol/l, potassium 5 mmol/l, chloride 117 mmol/l, bicarbonate 21 mmol/l, urea 5.5 mmol/l, creatinine 52 μmol/l, blood sugar 5.2 mmol/l and serum calcium was 2.2 mmol/l. Complete blood count was normal. Urine was negative for sugar, protein and abnormal sediments with an osmolality of 17 mmol/kg while that of the serum was 294. Urine volume passed over a 12-hour period was 700 ml. Administration of intranasal ADDVP did not result in any change in the biochemical values and osmolality of blood or urine. After an initial period of normal psychomotor development it was observed that her progress had begun to lag behind that expected for her age. At the age of seven years her psychomotor development was 12-months retarded and her weight and height were below the third centile for her age. Non-contrast computed tomography (CT) of the head was performed at that age (figures 1 and 2).

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**Questions**

1 What is the diagnosis?
2 What do the figures show?
Answers

QUESTION 1
Nephrogenic diabetes insipidus (NDI). The diagnosis is based on polyuria with a dilute urine, episodes of hypernatraemic dehydration and failure of intranasal ADDVP to increase urinary osmolality, the latter observation also rules out central diabetes insipidus.

NDI can be primary or secondary. Primary NDI usually affects males following an X-linked mode of transmission (MIM 304800); females are also affected to a lesser degree by an autosomal recessive pattern (MIM 222000). In this case, the parental consanguinity and the female sex favour an autosomal recessive inheritance. Secondary causes of diabetes insipidus (renal concentration defects) are shown in box 1. In this case secondary causes were ruled out.

QUESTION 2
Figure 1 shows bilateral symmetrical calcification of the basal ganglia and figure 2 reveals calcification at the junction between the white and grey matter in the parietal region.

Discussion
Causes of intracranial calcification in children are numerous (box 2). Intracranial calcification in children with primary NDI has been recently recognised and reported in a dozen cases. The cause(s) of intracranial calcification in patients with NDI is not exactly known. The postulated theory associates the intracranial calcification with hypernatraemic dehydration based on the following observations: recurrent episodes of hypernatraemic dehydration have occurred in all reported cases of NDI. Dehydration of this type is known to be associated with diffuse microscopic areas of intracranial haemorrhage and microthrombosis with tissue necrosis which subsequently calcifies. Also in favour of this theory is the finding of calcium deposits within and around blood vessels on brain autopsy of a patient who had NDI and intracranial calcification.

A direct relationship has been observed between the severity of intracranial calcification and the duration of the condition prior to diagnosis and establishment of treatment. This was noted in our patient where the calcification was mild, probably the result of less frequent attacks of dehydration. Mental retardation has varied from minor memory deficits to profound mental retardation and it has been recognised that the degree of psychomotor retardation is directly proportional to the degree of calcification. Growth retardation is frequent and probably related to caloric deprivation in infants and children who have to drink water in preference to food. Some believe, however, that growth retardation could be an inherent character of the disease.

Treatment of this condition requires a low sodium and protein diet, to decrease the osmotic load that the kidney has to excrete, in addition to a liberal fluid intake, particularly during febrile illness. The combination of thiazide diuretics and indomethacin therapy reduces the urine output dramatically. The mechanism of action of these drugs is not well understood. Diuretics increase the fraction of the filtered sodium chloride and water that is reabsorbed in the proximal tubule and as a result distal tubular flow is decreased. Indomethacin, a nonsteroidal anti-inflammatory agent, reduces renal prostaglandin production. This case clearly shows that primary NDI may be complicated by intracranial calcification accompanied by psychomotor retardation.

Causes of intracranial calcification in children
- inflammatory causes: cytomegalovirus infection, toxoplasmosis, neurocysticercosis, HIV infection, herpes simplex encephalitis
- vascular and hypoxic: hypernatraemic dehydration, hypoxic-ischaemic encephalopathy, calcified infarct, arteriovenous malformation
- tumours and dysplasia: calcified astrocytoma, neurenteric cysts
- endocrine disorders: hypoparathyroidism, hyperparathyroidism
- toxic disorders: carbon monoxide poisoning, lead poisoning, hypervitaminosis D, radiotherapy of leukaemia or tumours
- metabolic and hereditary diseases: mitochondrial encephalopathy, leukodystrophy, biotinidase deficiency, carbonic anhydrase II deficiency

Nephrogenic defects of urinary concentration

Conditions that affect primarily the action of antidiuretic hormone on tubular permeability to water
- hypokalaemia
- hypercalcaemia
- drugs (lithium, amphotericin B)
- hereditary (primary) NDI

Conditions that affect primarily the medullary solute concentration
- chronic renal failure
- obstructive nephropathy
- tubulointerstitial diseases (pyelonephritis, sickle cell disease, drugs)

Box 1

Box 2

Summary points
- primary NDI is transmitted as an X-linked recessive trait yet in a minority of cases it follows an autosomal recessive inheritance
- diagnosis depends on the passage of large volumes of diluted urine, episodes of hypernatraemia and unresponsiveness to administration of ADDVP
- this disorder is now recognised as being associated with intracranial calcification and brain damage. Early diagnosis and treatment may avoid such serious complications

Box 3
A high index of suspicion should be kept for this disorder, prompting early diagnosis and treatment in order to prevent brain damage.

**Final diagnosis**

Primary nephrogenic diabetes insipidus with intracranial calcification

**Keywords:** nephrogenic diabetes insipidus, hypernatraemic dehydration, intracranial calcification