Intra-atrial calcium infusions, growth, and development in end organ resistance to vitamin D

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Abstract
A five and a half year follow up of a girl with end organ resistant rickets is described. After failing to respond to high dose oral alfalcacidol (4 \( \mu \)g/kg/day) and calcium supplements, treatment for one year with domiciliary intra-atrial calcium infusions at 2 years of age induced a complete remission, which was maintained on subsequent high dose oral calcium supplement. Overnight infusions were well tolerated without adverse cardiovascular or renal sequelae or ectopic calcification. If the first three years of life are survived, the prognosis for a normal life on oral treatment is excellent.

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End organ resistance to 1,25 dihydroxyvitamin \( D_3 \) (1,25 \(( \text{OH})_2 D_3 \)) is a rare autosomal recessive disorder characterised by rickets which proves refractory to conventional vitamin D replacement. Cases have been reported from North America, North Africa, and the Middle East and it may be fatal if untreated in the first three years of life.\(^1\) DNA sequencing of the vitamin D receptor was established in 1988\(^2\) and the defect was shown to arise from point mutations of the ‘zinc fingers’ of the DNA binding domain of the vitamin D receptor later that year.\(^3\) Long term remission from rickets can be safely induced with an initial course of regular calcium infusions followed by oral treatment.

Case report
A girl with end organ resistance to 1,25 \(( \text{OH})_2 D_3 \) born in the UK to fourth generation Indian, non-consanguineous parents originating from Mauritius, has been followed up from the age of 4 months to 6 years. At 4 months of age she sustained a supracondylar fracture of the right femur and a retrospectively assessed skeletal survey demonstrated rickets. At 1·3 years she was referred to us with delayed motor milestones and was unable to weight bear. There was clinical, radiological, and biochemical evidence of severe rickets. Symptoms included irritability, apparent bone pain, anorexia, failure to thrive, weakness, motor and speech delay. Alopecia was striking (fig 1). Biochemical changes included hypocalcaemia, hypophosphataemia, secondary hyperparathyroidism, a raised alkaline phosphatase in excess of 10 000 IU/l and aminoacudria. A jejunal biopsy specimen, sweat test, and renal function measurement gave normal results. Two intramuscular injections of 300 000 IU of cholecalciferol (25(OH)D\(_3\)) at 1·3 and 1·4 years of age and daily oral alfalcacidol (1\(\alpha\)(OH)D\(_3\)) between 1·4 and 2·2 years up to a maximum of 30 \( \mu \)g/day, equivalent to 4 \( \mu \)g/kg/day, failed to induce a clinical response though the plasma 1,25 \(( \text{OH})_2 D_3 \) concentration was excessively raised at 400 pg/ml (normal range 30–50 pg/ml). Complete end organ resistance to 1,25 \(( \text{OH})_2 D_3 \) was subsequently confirmed on skin fibroblast cultures (A R Rut et al, personal communication). Normal receptor binding of 1,25 \(( \text{OH})_2 D_3 \) was shown but there was no in vitro conversion of 25 \(( \text{OH})_3 D_3 \) to 24,25 \(( \text{OH})_2 D_3 \).

INTRA-ATRIAL CALCIUM INFUSION REGIMEN
Intra-atrial infusions of calcium glubionate (calcium gluconogalactogluconate, Sandoz
Laboratories, Basle, Switzerland) were administered via a subcutaneous Port-a-Cath central venous access device with the catheter tip in the right atrium. Infusions began at an initial dose of 200 mg/day of the calcium salt made up to a volume of 500 ml with a solution of 5% dextrose, rising to 400 mg/500 ml/day on day 3, increasing to 500 mg/day from day 7 with nocturnal infusions from day 14. The parents learned the technique of aseptic infusions using a programmable pump and were supported by the nursing home care team.

![Graphs and charts](http://adc.bmj.com/)

**Figure 2.** Biochemical profile and growth against age before, during, and after calcium infusions. The period of calcium infusions is contained between the vertical dotted lines for all graphs. Normal ranges are indicated by the dashed horizontal lines. (A) Daily oral alfalcacidol (μg) and two intramuscular injections of cholecalciferol (300 000 IU). (B) Daily oral calcium and phosphate effervescent salts (mg) and daily intra-atrial (ia) calcium infusion dose (mg). (C) Plasma calcium (mmol/l). (D) Plasma alkaline phosphatase. (E) Plasma parathyroid hormone (ng/l). (F) Urinary calcium/creatinine ratio. (G) Patient height (cm). Note there were no episodes of hypercalcaemia during calcium infusions, but hypercalcaemia with oral calcium and alfalcacidol at the age of 5 (C). The parathyroid hormone fell to normal after three months of intravenous calcium infusions (E) but the alkaline phosphatase was slower to respond (D). The intra-atrial and oral doses of calcium are titrated to the interinfusion calcium/creatinine ratios (B, F). Growth crosses the 3rd centile during the calcium infusions, progressing to the 25th centile (G).
Calcium gluconate: 5% dextrose solutions were prepared by the hospital pharmacy weekly. Plasma concentrations of calcium, phosphate, urea, electrolytes, and creatinine, urinary calcium/creatinine ratios, and alkaline phosphatase activities were obtained by weekly and then monthly checks, as was height. Apart from one episode of the Port-a-Cath cloting soon after the start of the infusions, the system remained trouble free.

**CLINICAL AND BIOCHEMICAL COURSE DURING CALCIUM INFUSIONS**

Figure 1 illustrates the clinical picture before calcium infusions at 2-2 years. Figure 2 (A-G) plots the biochemical course and height before, during, and after the year of intra-atrial calcium infusions, indicated by the vertical dotted lines for all graphs. The infusions were well tolerated with no episodes of hypercalcaemia or electrolyte disturbance. Irritability was abolished and appetite restored within 24 hours of the start of the infusions. Attention, concentration, and language improved rapidly. Standing and walking with support began one week after starting infusions, and walking unsupported after four months of treatment at 2-5 years of age. Height achieved the 3rd centile for age after eight months (fig 2G). No cardiac arrhythmias or neurological sequelae were detected during the first two weeks of monitored infusions or subsequently. Renal function remained normal throughout and there was no evidence of nephrocalcinosis or ectopic calcification. Plasma concentrations of calcium and phosphate fluctuated at the lower limit of normal (2.1 and 1.0 mmol/l respectively) for the first month of infusions (fig 2C). The plasma immune parathyroid hormone concentration became normal after 12 weeks (fig 2E), falling from 249 ng/l to 31 ng/l (normal range 10–55 ng/l). Bone healing was complete by five months, though radiological change began 10 weeks after starting infusions (fig 3A, B), accompanied by a fall in alkaline phosphatase from 4325 IU/l to the upper limit of normal (fig 2D) and resolution of the aminoaciduria. Calcium infusions were titrated down to 120 mg/500 ml/12 hours against interinfusion urinary calcium/creatinine ratios (fig 2A, F).

**COURSE AFTER INFUSION**

Infection of the skin overlying the Port-a-Cath after one year required removal of the access device under antibiotic cover. Owing to a progressive rise in the parathyroid hormone and alkaline phosphatase after stopping calcium infusions (fig 2E, D), oral alfalcacidol, up to 20 μg/day, was restarted. This failed to prevent further rises, which only responded to the reintroduction of 1200 mg/day of oral calcium salt. At the age of 5, on 10 μg of alfalcacidol and 800 mg of oral calcium daily, the calcium/creatinine ratio (fig 2F) rose to 2.5 (normal <1), requiring further reductions in oral supplements (fig 2F). Growth continued along the 25th centile for height and there were no clinical symptoms of rickets.

**Discussion**

Reports of sporadic and multiple cases of 1,25 (OH)₂D₃ resistant rickets since the original description by Brooks in 1978 in a black American woman, have included North African or Middle Eastern families of Arabic descent in whom consanguinity is often, but not exclusively, a feature. The defect is characterised in our case by a point mutation in the amino acid sequence in the zinc finger region of the vitamin D receptor (A R Rut et al, personal communication) which normally binds to the nuclear DNA of the target cell for enzyme induction. There is normal 1,25 (OH)₂D₃ hormone to receptor binding. The
principal role of 1,25 (OH)2D3 is the active transport of calcium from the gut, though vitamin D receptors have been found in most tissues.7 However, pararenteral calcium alone may be sufficient for bone mineralisation.8 Fatal respiratory infections have been reported if the rickeys is not brought under control under the age of 3 years.1

As the graphs of the clinical and biochemical course of our case illustrate, long term calcium infusions appeared necessary to induce a remission from total body calcium depletion, which became clinically and radiologically gross by the age of 4 months due to exhaustion of placentally acquired calcium stores. Our data suggest that oral calcium supplements appeared to induce a slow biochemical change, which may have been independent of a high dose alfalcidol intake of up to 30 μg/day before calcium infusions. This was not matched by clinical or radiological improvement. Intravenous calcium infusions resulted in a rapid clinical, radiological, and biochemical recovery similar to short term results that had been reported in three previous case series involving four patients5 6 9 by the time we began infusions in March 1989. Unlike a recently published report of long term follow up in 10 cases spanning two kindreds,10 biochemical resolution in our case was complete after three months of infusions.

We did not have problems of hypercalcaemia and bradycardia as described by Hochberg et al.10 In contrast to the mean septicaemia onset time of eight months in the Hochberg series, our patient never became septicaemic but developed an infection around the Port-a-Cath after one year of needle punctures. A subcutaneous implant may protect against sepsis as well as offer greater freedom for normal activities. Continued improvement on oral calcium supplements remains intriguing as little calcium is absorbed enteraly in the absence of functioning vitamin D receptors.11 Low plasma calcium concentrations normally induce conversion of 25 to 1,25 (OH)2D3 which stimulates active transport of calcium from the gut. Conversely, enteral calcium absorption is down regulated via the concentration of 1,25 (OH)2D3. Once bones are calcium replete, high dose oral calcium maintains remission.

CONCLUSION
Monotherapy with domiciliary calcium infusions for one year induced a striking remission of severe rickets in a 2 year old girl who had proved refractory to initial oral treatment. Infusions with appropriate outpatient and home care nursing back-up, proved practical and safe. A urinary calcium/creatinine ratio of >1 between infusions indicates the need to reduce the daily calcium intake, and a normal parathyroid hormone is the best guide to continued remission, whether calcium is administered parenterally or orally.

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