Vitamin D has steroid hormonal effects which can produce clinical symptoms and signs unrelated to calcium homeostasis. Its deficiency has been implicated as a risk factor for diabetes, ischaemic heart disease, and tuberculosis in Asians. In this review, the incidence, aetiology, prevention, and treatment of symptomatic vitamin D deficiency in childhood are considered. A renewed public health campaign is required in the UK to address the continuing problem of vitamin D deficiency in Asian families.

Vitamin D has wide ranging steroid hormonal effects which can produce clinical symptoms and signs unrelated to calcium homeostasis. The explanation appears to be that the receptor for 1,25 vitamin D is present in most cells of the body, and there is widespread extrarenal expression of 1-α hydroxylase, which catalyses synthesis of this hormone. 1,25 Vitamin D has a regulatory role in skeletal muscle, the immune system, and cell differentiation, including induction of proteins, such as nerve growth factor, in the central nervous system. Recently vitamin D deficiency has been implicated as a risk factor for diabetes, ischaemic heart disease, and tuberculosis in Asians.

Prenatal famine and adult disease is now being increasingly recognised. Low maternal vitamin D may adversely affect the developing fetal brain. In addition to the previously known paediatric problems of hypocalcaemic fits, dental enamel hypoplasia, infantile rickets, and congenital cataracts in early life, vitamin D deficiency has been shown to affect postnatal head and linear growth.

In the UK, the Committee on Medical Aspects of Food Policy (COMA) recommends that all infants receive 7–8.5 µg (280–340 iu) of vitamin D daily either in a multivitamin preparation or in fortified infant formula milk. It recommends that all pregnant and lactating mothers should receive 10 µg (400 iu) of vitamin D daily, and advises that Asian children should be encouraged to take vitamin D supplements throughout the first five years of life.

CHANGING INCIDENCE

Deprivational vitamin D deficiency in Asians living in the UK, whose origins are from India, Pakistan, or Bangladesh, was first reported nearly 30 years ago. Various campaigns to improve life and social conditions, as well as recommendations advocating provision of free vitamin D supplements led to a decline in the prevalence of vitamin D deficiency in these ethnic groups.

A questionnaire survey, commissioned by COMA in 1995, among West Midlands paediatricians, confirmed that the majority of the respondents reported seeing very few cases of rickets and when seen, were predominantly in Afro-Caribbean toddlers rather than Asian infants. However, we have seen a resurgence of vitamin D deficiency in Asian children in the past two to three years in Birmingham, experience mirrored by reports from elsewhere in the UK of vitamin D deficiency in infants (London), adults (Leicester), and pregnant women (Cardiff). In addition to rickets, infants have presented in the first six to nine months of life with symptomatic hypocalcaemia, particularly seizures; we have seen one case of life threatening cardiomyopathy.

Unlike other reports which have implicated prolonged and exclusive breast feeding, most of our infants have been both breast and formula fed and several have been receiving multivitamin supplements. These infants have all had evidence of severe vitamin D deficiency (25-hydroxyvitamin D <8 ng/ml) as have the majority of the mothers who have also been tested. Most of these infants have been born to mothers who were themselves born in the UK. It would appear that the previous recommendations regarding prevention have been largely forgotten and our initial enquiries indicate that most UK maternity units do not have a policy of providing vitamin D supplements to pregnant Asian women.

AETIOLOGY

The presentation of our cases and others highlights the importance of maternal vitamin D stores. Newborn infants depend on fetal stores of vitamin D obtained from their mother; following delivery their vitamin D status is 60–70% of measured maternal vitamin D concentrations.

No meta-analysis has been performed to assess the outcome of maternal vitamin D deficiency as the populations studied are either too diverse or contain too small numbers with the reports failing to provide summary statistics. Ongoing reports from throughout the world over a 20 year period show that a causal relation probably does exist between clinical symptoms and infant and maternal vitamin D deficiency.

A study of 25-hydroxyvitamin D concentrations in the resident non-pregnant adult population in the catchment area of one Birmingham hospital, showed that in winter, 85% of Asians, compared to 3.3% of non-Asians, had vitamin D concentrations less than 8 ng/ml. Furthermore during summer, 38% of the Asians still had
vitamin D concentrations less than 10 ng/ml. Previous reports from the UK and other European countries have also found differences between Asian and non-Asian groups. Shaunak and colleagues found 22% of healthy adult Asian couples (both sexes) had vitamin D deficiency (25-hydroxyvitamin D < 10 ng/ml) with normal concentrations in white controls. A recent survey in Cardiff of vitamin D concentrations in pregnant women showed that 54% of Asian women had biochemical vitamin D deficiency (< 8 ng/ml). A survey undertaken in Norway showed that 83% of pregnant Pakistani women were vitamin D deficient.

Previous studies have attributed vitamin D deficiency in Asian women to cultural and dietary habits and their infrequent use of supplementation. The main source of vitamin D is from ultraviolet irradiation of the skin; there is no radiation of the appropriate wavelength (290–310 nm) in Britain from the end of October to the end of March. For the remaining months of the year, 60% of the effective ultraviolet radiation occurs between 1100 and 1500. Although lack of sunlight exposure is important, this is probably not the only factor, as the vitamin D study undertaken in Birmingham also showed a high prevalence in men. Furthermore, a recent study from Delhi has found that excessive numbers of healthy Asian subjects have low 25-hydroxyvitamin D concentrations while residing in abundant sunlight.

In order to answer this dilemma, work undertaken in the USA has shown that altered vitamin D metabolism caused by notably increased 25-OHVD24 hydroxylase activity may be responsible for low 25-hydroxyvitamin D concentrations in Asians. Thus, it appears they may have a genetic predisposition to vitamin D deficiency. This suboptimal vitamin D status facilitates the occurrence of symptomatic vitamin D deficiency when there are increased requirements, as in early childhood, adolescence, pregnancy, and lactation.

Normally the calcium demands of pregnancy and lactation occur through a two- to threefold upregulation of intestinal calcium absorption in late pregnancy and during lactation. This is mediated by a similar twofold pregnancy induced rise in 1,25-dihydroxyvitamin D, during the second and third trimesters with a concomitant reciprocal 50% fall in parathormone. We postulate that in vitamin D deficient Asian women during pregnancy, the mechanism of calcium upregulation by a rise in 1,25-dihydroxyvitamin D is lacking and hence vitamin D resources are restricted to the fetus.

**PREVENTION**

It is apparent that relying on vitamin D supplements given to infants or the amounts present in formula feeds is inadequate to overcome the impact of maternal vitamin D deficiency. It is important to ensure that fetal stores are optimised by supplementation of at risk mothers. A 400 iu daily dose of vitamin D should theoretically result in normal concentrations of 25-hydroxyvitamin D, and 1,25-dihydroxyvitamin D. However, a recent study from Denmark undertaken in veiled Moslem women indicates that, in the absence of sunlight exposure, a dietary intake of 600 iu vitamin D per day is insufficient to maintain adequate 25-hydroxyvitamin D concentrations; the authors suggested that 1000 iu per day would be more effective. Screening pregnant Asian women for vitamin D status and supplementing as necessary has been suggested. However, this can only be advocated after clarifying a nationally agreed "deficient range" for vitamin D, the optimal time to screen in pregnancy, and the most appropriate vitamin D dose to achieve compliance, efficacy, and minimal risk.

It would be simpler to supplement all pregnant Asian women with vitamin D. There are, however, problems in compliance with vitamin D supplementation, particularly when intended to be taken daily. A recent health programme in Norway, consisting of free samples of 10 µg (400 iu) vitamin D daily combined with information, failed to reduce vitamin D deficiency to acceptable low concentrations in Pakistani mothers. An alternative is to give 25 µg (1000 iu) daily during the third trimester, which has been shown to produce normal 25-hydroxyvitamin D concentrations in mothers and infants at term. Another proposal is a single high dose of vitamin D of 100 000–200 000 iu given during the sixth or seventh month of pregnancy, which appears sufficient to cover both maternal and fetal needs. Adequately controlled antenatal vitamin D supplementation trials are few, although Brooke et al in London and Mallet et al in northwest France, who performed case matched cohort studies, identified that antenatal maternal vitamin D concentrations are correlated with infant status.

As regards infants, it is clear that national recommendations about vitamin D supplements are not being followed. We believe all Asian infants should be given 400 iu vitamin D daily whether breast or formula fed. Alternatively, if poor daily compliance is a problem, an annual dose (4 mg, 150 000 iu) at the beginning of autumn, appears to provide protection against vitamin D deficiency without vitamin D overload.

**TREATMENT OF SYMPTOMATIC VITAMIN D DEFICIENCY IN CHILDHOOD**

The most effective and physiological treatment for an infant with symptomatic hypocalcaemia, or rickets caused by vitamin D deficiency, is either ergocalciferol (vitamin D2) or cholecalciferol (vitamin D3). This is the fastest means to replenish the depleted 25-hydroxyvitamin D concentrations, improve intestinal calcium absorption, and suppress the effect of secondary hyperparathyroidism. Alfacalcidol or calcitriol are not appropriate treatments, and children treated with these preparations initially are often resistant to their effect. In the UK, preparations of ergocalciferol or cholecalciferol are not readily available and a liquid preparation suitable for infants is not listed in the British National Formulary (BNF). A calciferol solution containing 3000 units per ml is available from Martindale Pharmaceuticals (Bampton Road, Harold Hill, Romford, Essex RM3 8UG) and is listed in Medicines for Children. A dose of 3000 units daily for infants less than 6 months and 6000 units daily for those older than 6 months for six weeks to four months is usually adequate to replenish depleted vitamin D stores, correct biochemical abnormalities, and heal rickets. Oral calcium supplements are also usually required initially if the child is hypocalcaemic and/or dietary intake is poor. It is not necessary to continue treatment until any leg deformity resolves, which may take up to two years. For the occasional adolescent presenting with symptomatic hypocalcaemia, a tablet preparation containing 10 000 units of cholecalciferol is listed in the BNF and is the most appropriate for this age group. In addition to monitoring calcium, phosphate, and alkaline phosphatase while on treatment, serum parathormone is useful as it should normalise when the vitamin D deficiency has resolved.

**RECOMMENDATIONS**

It is our opinion that a renewed public health campaign is required in the UK to address the continuing problem of vitamin D deficiency in Asian families. Local circumstances will probably dictate what are the most cost-effective measures. Our current recommendations would be:

- The definition of 25-hydroxyvitamin D deficiency should be standardised in the UK to < 8 ng/ml during winter and < 10 ng/ml during summer, so that reports of prevalence can be compared.

- All pregnant Asian women should receive vitamin D supplements—either 400 iu daily from the booking clinic appointment or 1000 units daily during the third trimester. Alternatively a single dose of ergocalciferol (100 000–
Vitamin D deficiency in UK Asian families

200 000 iu) during the sixth or seventh month may be preferred to achieve best efficacy and compliance.

- All Asian infants, whether formula or breast fed, should receive 400 iu vitamin D daily. Alternatively an annual regimen may provide a higher compliance rate.

- Awareness of the risks of vitamin D deficiency should be increased in all Asian families.

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