Dysequilibrium/ataxic diplegia with immunodeficiency

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Abstract
A girl with purine nucleoside phosphorylase (PNP) deficiency is described. The nature of the motor disorder is similar to other children since found to have PNP deficiency. It is suggested that the diagnosis be considered in any child with unexplained dysequilibrium/ataxic diplegia. Other previously unreported features are intracytoplasmic neutrophil inclusion bodies and an improvement in the neutropenia after intravenous immunoglobulin.

Severe congenital immunodeficiency states generally present with recurrent infections as maternal passive immunity declines. Children with purine nucleoside phosphorylase (PNP) deficiency can, however, present with a non-progressive motor disorder before the onset of the immunodeficiency giving an opportunity for treatment before secondary complications occur and for genetic advice. This requires a clearer definition of the nature of the motor disorder than had previously been available.

Case history
A 1 year old girl presented with delay in motor development. She had been rolling since 9 months, could sit supported only with a ‘C’ shaped back, and could not crawl or weight bear. She gave the appearance of hypotonia, although examination demonstrated slightly increased tone and brisk reflexes. She could use her hands crudely, tucking in her thumbs, but had a full range of supination. She appeared bright and alert. Birth had been normal with a birth weight of 4400 g. Her parents were healthy and unrelated. Computed tomography of the brain was normal. She learned to sit, propping herself, at 16 months and at 19 months was able to pull herself to standing. Although her balance remained very poor, she was able to walk with her hands held at 26 months and at 3 years take a few unsteady steps. Fine hand movements were mildly static, her speech clear, and intellectual development normal.

At age 26 months she began to suffer severe recurrent urinary tract infections despite prophylactic antibiotic treatment. She had had no previous significant infections. Ultrasonography showed pelvicalical dilatation, progressing to frank hydronephrosis within six months. A micturating cystogram confirmed severe vesicoureteric reflux. Haematological indices were normal at 33 months.

When admitted with a further urinary tract infection at 3 years she was noted to have oral candida, right cervical lymphadenopathy, and to be both neutropenic (neutrophil count 1.0×10⁹/l and lymphopenic (lymphocyte count 1.2×10⁹/L). A bone marrow aspirate showed megaloblastoid nuclei, dyserythropoiesis, and myeloid maturation arrest with bizarre intracytoplasmic inclusion bodies; this was also seen in peripheral blood neutrophils (figure). Immunoglobulins were normal, tetanus toxoid antibodies were present in normal amounts, isohaemagglutinins were weakly present, nitroblue tetrazolium test was equivocal, and HIV antibody was negative. Total lymphocyte and T₄ and T₃ subset numbers were severely depressed (0.4×10⁹/l, 0×10⁹/l, and 0.02×10⁹/l respectively) and were functionally unresponsive to mitogens. Adenosine deaminase activities were raised, however, and PNP was absent. Maternal and paternal values were 53% and 58% of normal respectively. A lymph node biopsy specimen showed lymphoblastic/imnunoblastic non-Hodgkin’s lymphoma.

Her clinical condition deteriorated and as there was no compatible bone marrow donor it was decided, after discussion, to give purely palliative lymphoma treatment. The child died 20 days later.

Discussion
PNP deficiency was first described in 1975. Enzyme substrates inosine, deoxynosine, guanosine, and deoxyguanosine accumulate and enzyme products urate and guanosine triphosphate are depleted. The principle systems affected are the lymphoid system and central nervous system.

Neurological dysfunction primarily affecting
motor development is clearly recognised in association with PNP deficiency, often being the presenting feature. The characteristics of the motor disorder have not been clearly defined, however, with previous papers referring to spastic quadriplegia or spastic diplegia. Two siblings described as having familial dysequilibrium/diplegia with T lymphocyte deficiency have now retrospectively been found to have PNP deficiency, using cultured fibroblasts, at this hospital. Both parents had heterozygote values for PNP. These children were thought to be similar to those described by Hagberg et al as having familial ataxic diplegia with deficient cellular immunity. The four children described in these two papers, plus our own case, point to a fairly characteristic and identifiable clinical picture. This would seem to consist of a disequilibrium—that is, pronounced difficulty in maintaining posture and upright position giving rise to an early hypotonia, very unsteady and delayed sitting, and great difficulty in walking independently. In contrast there is either no, or only a mild, intention tremor or clumsiness. There is evidence of a spastic diplegia on clinical examination with brisk reflexes, upgoing plantars, and sometimes ankle clonus and restriction of passive dorsiflexion of the ankles. When walking this is masked by trunkal ataxia, however, particularly as the child gets older. The neurological picture appears to be non-progressive, may relate to the degree of guanosine triphosphate depletion and is non-infective in aetiology. Intelligence may be normal as in this case.

The haematological and immunological consequences of PNP deficiency relate to the depressed T cell numbers and function, leading to repeated infections with viral illnesses often being the cause of death. These children should not receive live vaccinations. Lymphomas, often of a B cell origin, are also a complication of other immunodeficiency states, for example, AIDS, and have been described in PNP deficiency. Haematological findings in PNP deficiency are pure red cell aplasia, megaloblastic anaemia, autoimmune haemolytic anaemia, and maturational arrest at the myelocyte/metamyelocyte stage in bone marrow aspirates. Intracytoplasmic neutrophil inclusions (figure) have not been previously described; they persisted after resolution of sepsis and in addition, identical inclusions were seen in the paternal blood film. Similar inclusions are well recognised during treatment with drugs that interfere with DNA synthesis such as azathioprine, chlorambucil, and appear to be fragments of DNA. DNA synthesis is impaired in PNP deficiency suggesting a similar aetiology.

Autoimmune disease is another complication of immune deficiency including PNP deficiency. Suspecting the neutropenia to be immune mediated, we treated our patient with 2 g/kg intravenous immunoglobulin over two days. There followed a dramatic rise in the neutrophil count (0.4–2.8 x 10^9/l). The patient had been persistently neutropenic for two months before this. In conclusion, recognition of PNP deficiency before onset of severe infection or malignancy is important so that bone marrow transplantation from a suitable donor can be considered while the patient is well. Successful engraftment has already been documented in this condition. In addition, antenatal diagnosis is now possible for measuring PNP in chorionic villus samples. We consider familial dysequilibrium/diplegia with T lymphocyte deficiency and familial ataxic diplegia with deficient cellular immunity should no longer be considered separate disease entities, but reclassified as cases of PNP deficiency.

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Fulminant hepatitis B in an infant born to a hepatitis B antibody positive, DNA negative carrier

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Abstract

A boy was born to a mother who was a chronic hepatitis B virus (HBV) carrier. She was hepatitis B (HBe) antibody positive and HBe antigen and HBV-DNA negative. The boy had not received hepatitis B vaccine and died from fulminant hepatitis at 3 months of age. This case demonstrates the need to vaccinate babies of HBe antibody positive, HBe antigen negative carriers.