Case Report

An Unknown Autoinflammatory Syndrome Associated with Short Stature and Dysmorphic Features in a Young Boy

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ABSTRACT. A young boy from nonconsanguineous Palestinian parents presented with short stature, motor development delay, wide nasal bridge, bilateral periorbital edema, everted lower lip, brachydactyly, large interphalangeal articulations, drumstick extremities of the fingers, bilateral simian crease, clindactyly of the 5th fingers, painful joints, subcutaneous nodules all over his body and recurrent episodes of fever of unknown origin. Differential diagnoses such as the hyperimmunoglobulinemia D syndrome, tumor necrosis factor receptor associated periodic syndrome (TRAPS), the chronic infantile neurological cutaneous and articular (CINCA) syndrome, and the newly recognized nodulosis, arthropathy, and osteolysis (NAO) syndrome are discussed. This syndrome may not have been previously reported. (J Rheumatol 2002;29:1084–7)

Key Indexing Terms:
FEVER OF UNKNOWN ORIGIN
SUBCUTANEOUS NODULES
PERIORBITAL EDEMA
PAINFUL JOINTS

Autoinflammatory syndromes are systemic disorders characterized by apparently unprovoked inflammation in the absence of high titer autoantibodies or antigen-specific T lymphocytes1. The hyperimmunoglobulinemia D syndrome (HIDS), familial Mediterranean fever, systemic onset juvenile rheumatoid arthritis, the nodulosis, arthropathy and osteolysis (NAO) syndrome, the tumor necrosis factor (TNF) receptor associated periodic syndrome (TRAPS), and the chronic infantile neurological cutaneous and articular (CINCA) syndrome are part of this group.

We describe a patient with short stature, dysmorphic features, brachydactyly, large interphalangeal articulations, and unexplained episodes of fever with inflammation consisting of periorbital edema, painful joints, and subcutaneous nodules.

CASE REPORT
The boy described here is the son of healthy nonconsanguineous parents who are Palestinian refugees in Lebanon. The boy has 3 healthy brothers and sisters. Three other sisters died at less than one week of age from unknown reasons. Medical followup during the pregnancy was not performed, but the parents did not recall any toxic exposures or unusual events during that period. Delivery was at 32 weeks of amenorrhea. At that time, the mother’s age was 17 years, and the father’s 18. Birth weight, length, and head circumference were not reported. At birth, the boy had a large nasal bridge, large interphalangeal articulations, and subcutaneous nodules all over his body (Figure 1). Since birth, he has had recurrent febrile attacks with fever as high as 39–40°C for 3–4 days without chills. Those episodes were accompanied by painful joints and periorbital edema, and followed by weakness lasting about a week. Results of investigations performed several times following those attacks were normal. Developmental delay was noted during the first years of life, as the boy was not able to walk without help until the age of 4 years.

At the time of report, the boy was 10 years old. His intellectual development seemed normal. On examination, his weight was 21.3 kg (< 3rd centile), height 118 cm (<< 3rd centile), and head circumference (OFC) 51 cm (10th centile). He presented with bilateral periorbital edema, wide nasal bridge, especially in its middle part, everted lower lip (edematous on the right side), bifid uvula, brachydactyly (hand length 13.5 cm: ~ 3rd centile), large interphalangeal articulations, drumstick extremities of the fingers, bilateral single palmar creases, clinodactyly of 5th fingers, and flat feet. Joints were painful but without limitation of articulations. Subcutaneous nodules were found all over his body, especially over the limbs, along with a few dark red cutaneous spots (Figure 2). No rashes were noted. The external genitalia were normal. Neurological examination was unremarkable, except for the presence of a mild waddling gait. Deep tendon and abdominal reflexes were present. Plantar responses were in flexion. Ophthalmological evaluation was normal. ENT examination disclosed a perforation of the left eardrum.

Skin specimens were obtained for biopsy and analyzed by 2 different pathologists. Both reported a leukocytoclastic vasculitis. Analysis of 2 subcutaneous nodules showed chronic lymphadenitis. The radiological examination of the skeleton was unremarkable except for the presence of a...
slight external deviation of the last phalanges of both 2nd fingers. A magnetic resonance image of the brain showed no gross malformations.

Abdominal ultrasounds, cardiac echography, and the auditory brainstem response were all normal.

Complete blood count, hemoglobin electrophoresis, blood glucose levels, urinalysis, amino acid assay of plasma and urine, urinary excretion for mucopolysaccharidosis, ceraminidase and mevalonic acid excretion in urine, liver and thyroid function studies, CPK, aldolase, LDH, and VDRL serology were all generally within normal levels. Tests for nDNA, RNP, Sm, SSA, SSB, SC1-70, JO1, and antiplatelet antibodies were negative.

Antinuclear antibody result was positive. Elevated serum C-reactive protein levels were found. Chromosome study of lymphocytes with high resolution in G and R banding showed a normal 46,XY karyotype.

Mutations in the marenostrin/pyrin gene for familial Mediterranean fever were investigated by fluorescent sequencing of exon 10 and by polymerase chain reaction digestion for the E148Q mutation in exon 2. No mutation was identified.

DISCUSSION

The main phenotypic manifestations of this patient consisted of short stature, dysmorphic features, brachydactyly, large interphalangeal articulations, and unexplained episodes of fever with inflammation consisting of peri orbital edema, painful joints, and subcutaneous nodules. The disease started to manifest itself directly after birth, but had not caused any detectable psychomotor deterioration by age 10. Differential diagnoses were discussed that included autoinflammatory syndromes and syndromes or diseases that present with subcutaneous nodules and/or peri orbital edema.

The hyperimmunoglobulinemia D syndrome is characterized by early onset of attacks of periodic fever and elevated serum polyclonal IgD (> 100 U/ml). Symptoms during attacks include joint inflammation, skin lesions, swollen lymph nodes, headache, and abdominal complaints. In our patient, the normal value of IgD and urinary mevalonic acid allowed us to rule out this syndrome. Chronic infantile neurological cutaneous and articular syndrome is a chronic inflammatory illness that most often starts at birth and persists throughout life. It is characterized by a triad including arthropathy, cutaneous rash, and chronic meningitis. Arthropathy was present in our patient, but the 2 other features of the triad were absent, allowing a simple differentiation. Finally, TNF receptor associated periodic syndrome, an inflammatory disorder characterized by prolonged episodes of periodic fever, localized inflammation, and a spectrum of dermatologic findings including migratory patches, edematous plaques, peri orbital edema and/or conjunctivitis, was not retained as a possible diagnosis as our patient never presented skin eruption or abdominal pain, but had short stature and dysmorphic features not reported in TRAPS. Also, skin biopsy specimen showed leukocytoclastic vasculitis but no perivascular dermal infiltrate of lymphocytes and monocytes, as observed in TRAPS.

Systemic onset juvenile rheumatoid arthritis, juvenile hyaline fibromatosis, and Henoch-Schönlein disease may present with subcutaneous nodules, fever, painful joints, and a variety of other extraarticular features. They were ruled out as our patient showed no signs of arthritis, joint contractions, nodular skin lesions, generalized osteopenia, or any other signs typically associated with those entities. The newly recognized nodulosis, arthropathy, and osteolysis (NAO) syndrome was ruled out as well, since our patient did not have osteolysis with carpal and tarsal resorption and osteoporosis, and because he had subcutaneous nodules all over his body, whereas it is confined to the palmar and plantar region in NAO syndrome.

Farber disease (MIM 228000), a rare condition that may present with painful joints, subcutaneous nodules, and intermittent fevers, was ruled out because of the normal value of the acid ceramidase enzyme.

Peri orbital edema may occur initially or in the course of a wide variety of diseases such as hypo- or hyperthyroidism, hypoalbuminemia, angioedema, trichinosis, allergic reactions, neoplastic diseases, sarcoidosis, dermatomyositis, systemic lupus erythematosus (SLE), and lupus panaritulitis (LP). From all these, only the last 3 were candidates for...
Figure 2. A–D. The patient at age 10 years. Note the bilateral periorbital edema, wide nasal bridge, dark cutaneous spots, and edematous everted lower lip.
differential diagnosis, all the more because subcutaneous nodules and periorbital edema have been reported in SLE and LP, sometimes as the first clinical feature of the disease\cite{8,9}. Nevertheless, these diseases were ruled out as neither the clinical description nor the pathological result in our patient was consistent with one of them.

This patient presented an inflammatory syndrome with a combination of anatomical features never reported before. It is hoped that this report will alert physicians to the existence of this combination and lead to additional reports confirming that it is a new entity.

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REFERENCES