Panniculitis: a report of four cases and literature review

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Abstract
Panniculitis is a disease with many causes and associations. The classification of clinical subtypes is unsatisfactory and hampered by the use of eponyms.

Four children with recurring panniculitis are described and their histology presented. Three had subcutaneous fat atrophy with lobular panniculitis on biopsy; all responded well to corticosteroids. The fourth child had a septal panniculitis with no atrophy of subcutaneous tissues and only a partial response to treatment with corticosteroids.

A widely accepted precise histological classification of panniculitis is needed to enable accurate predictions of the outcome of this serious disorder.

Panniculitis is an uncommon but fascinating condition in childhood. Inflammation of the fat may arise spontaneously or occur as a feature of distinct illnesses such as pancreatitis,1 bacterial sepsicaemia,2 systemic lupus erythematosis,3 polyarteritis,1 and malignancy.4 The literature abounds with isolated case reports using eponyms such as 'Weber–Christian syndrome'5 and 'Rothman–Makai panniculitis'.6 Although attempts have been made to classify histological subtypes of panniculitis,1,7 no universally recognised classification is in use to aid retrospective studies. The various forms of lipodystrophy are often included in the classification of panniculitis,8 which then becomes further confused.

We report four cases of spontaneously occurring panniculitis, one in an infant and three in older children. Follow up of our four children over several years suggests that lobular panniculitis is associated with significant fat atrophy at the site of previously active lesions and a good response to corticosteroids. Septal panniculitis, however, has no associated atrophy of fat and only a partial response to corticosteroids.

Case reports
CASE 1
A 6 year old girl first presented in 1975 with an upper respiratory infection and painful lumps on both thighs. Both problems resolved spontaneously. The painful lumps reappeared three years later and biopsy specimen of a nodule demonstrated lobular panniculitis (fig 1).

Other investigations included a haemoglobin concentration of 94 g/l, total white cell count of 11·9×10⁹/l, erythrocyte sedimentation rate (ESR) of 60 mm/hour, a significant rise in antistreptolysin O titre (ASOT) over two months, a positive rheumatoid factor, and raised immune complexes. Serum C3 concentration was 1·65 g/l (normal range 0·8–1·70 g/l) and C4 0·23 g/l (normal range 0·15–0·45 g/l). Serum IgG was 28 g/l (normal range 6·3–16·5 g/l). Direct Coombs test, the Venereal Disease Research Laboratory test, Treponema pallidum haemagglutination assay, and tests for anti-nuclear and anti-DNA antibodies gave negative results. No cryoglobulins or cold agglutinins were detected, and serum IgA, IgM, euglobulin lysis and fibrin plate lysis were normal.

Prednisolone was given in a dose of 30 mg/day orally from January 1979 tapering to 30 mg and 5 mg on alternate days by February 1979. The prednisolone dose was steadily reduced over the next five months, ceasing in July 1979 at which time there was resolution of the lesions with appreciable loss of subcutaneous fat at the sites of the previous nodules. The nodules reappeared three months after cessation of treatment. Treatment with prednisolone was again successful in controlling the symptoms when given in a dose of 30 mg/day orally, initially, and decreased steadily as before over the next nine months until ceasing in July 1982. Currently the child remains well and free of new lesions.

CASE 2
This girl was completely normal until the age of 5 months when she developed red hot lumps on both legs. By 16 months of age the lesions had...
spread to involve her face, arms, and trunk. There were no systemic symptoms. A skin biopsy specimen showed a septal panniculitis (fig 2) that responded temporarily to prednisolone in small doses.

At 3-5 years of age she developed fever, joint pains, and stiffness with increasing numbers of nodules appearing after bouts of tonsillitis. A trial of prophylactic penicillin decreased the number of infections but did not appear to modify the disease process.

Subsequent treatment included chloroquine, indomethacin, naproxen, and sulphasalazine; none of these drugs resulted in any lasting benefit. Prednisolone was used in small doses (for example, 5 mg) daily until the age of 5-5 years when alternate day treatment was begun up to a dose of 20 mg at 11 years old. Increases in prednisolone dose led to partial improvements but relapses still occurred (fig 3). Temporary improvement was seen after three pulses of intravenous methylprednisolone were given in response to an acute exacerbation of the disease. Dapsone was started at 13 years of age with some improvement of fevers and nodules.

During the course of this girl's illness her haemoglobin has remained at 100-111 g/l and her ESR has ranged from 13 to 30 mm/hour.

Tests for antinuclear antibody, immune complexes, autoantibodies to gastric parietal cells, smooth muscle, mitochondria, reticulin and thyroglobulin gave negative results. ASOT, throat swab, and urine and blood cultures were also negative. Plasma urea, electrolytes, C reactive protein, triglycerides, cholesterol, red cell folate, serum complement, immunoglobulins, and B12 concentrations and bone marrow examination were all normal.

**CASE 3**

In December 1985 an 11 year old girl developed painful erythematous indurated lesions over her feet and legs (fig 4) after recovering from a sore throat. She was not febrile but appeared lethargic. At the same time the metatarsophalangeal joints of both feet were tender.

There was no improvement with penicillin, aspirin, other non-steroidal anti-inflammatory drugs, and bed rest. Her haemoglobin concentration was 113 g/l, white cell count 6.8 x 10⁹/l, and ESR 60 mm/hour. Rheumatoid factor was positive (RAHA 1:1280) initially, becoming negative by November 1987. Plasma C reactive protein was 27 000 µg/l (normal <8000). The
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plasma urea and electrolytes, liver function tests, amylase activity, ASOT, serum immunoglobulins and complement, urine analysis, nose and throat swabs, and chest radiography all gave normal results. Autoantibodies to parietal cells, mitochondria, smooth muscle, reticulin, thyroid microsomes, and thyroglobulin were not detected. Antinuclear and anti-DNA antibodies were negative and antibodies to extractable nuclear antigen were not detected.

One month after onset of the illness nodules began to appear on the child's arms. A skin biopsy specimen demonstrated lobular panniculitis and prednisolone was started at a dose of 40 mg alternating with 10 mg daily. She became more energetic and the skin lesions gradually resolved, leaving large areas of subcutaneous fat loss (fig 5). Prednisolone intake was cautiously reduced as three small lesions persisted and occasional lesions appeared at sites of minor skin trauma. By August 1988 the prednisolone dose was 25 mg on alternate days, the child felt well, and the ESR had fallen to 6 mm/hour.

CASE 4
A girl weighing 4500 g was born by elective caesarean section after a normal pregnancy in August 1985. She had signs of ventricular and atrial septal defects leading to congestive cardiac failure by 5 weeks of age. Corrective cardiac surgery was performed three weeks later after medical treatment had failed to control her condition.

Postoperatively the baby required intensive respiratory and cardiovascular support but was able to go home after two weeks in hospital. Shortly afterwards she developed red tender lumps on her trunk, thighs, arms, and scalp. These lumps resolved spontaneously but reappeared five weeks later on the baby's arms, shoulder, and face where they were sufficiently tender to interfere with feeding.

A biopsy specimen of a active lesion showed lobular panniculitis that responded well to prednisolone in a dose of 1-5 mg/kg/day. The dose of steroid was reduced over the next five months and there has been no recurrence of disease off treatment.

Investigations at the time of biopsy showed a haemoglobin concentration of 95 g/l, white cell count of 12.2×10⁹/l, and an ESR of 13 mm/hour. Serum amylase and creatine phosphokinase activities and ASOT were normal. No immune complexes were detected.

Discussion
Our four cases demonstrate the two histological types of panniculitis previously described, lobular and septal. Although these types have been recognised in the past no attempt has been made to correlate histological type with response to treatment. Our review of the literature supports this paper's finding that lobular panniculitis responds better to treatment with prednisolone than the septal type.

Two of our infants were infants at the onset of the illness. Most of the infants described previously have had significant systemic symptoms and in some cases a fatal outcome. Our infant with lobular panniculitis has done well on steroids alone. She now has no fat atrophy and is off treatment with no recurrences. Taylor and more recently Conway et al10 have also described cases of panniculitis with onset in infancy where the clinical course has been relatively mild and there has been a good response to treatment. Our child with septal panniculitis whose disease began at 5 months of age, although still having problems, has not succumbed and has not developed symptoms of perivisceral disease.

The nature of the inflammatory infiltrate found on histology varies according to the disease stage at which the biopsy is performed. 11 It is important to obtain adequate biopsy specimens early in the disease as once the end stage is reached, subcutaneous fibrosis of the panniculus may be all that is seen and the lobular architecture of the fat is distorted. Punch biopsies are not recommended because they may not obtain adequate samples of fatty tissue because of inflammation of the subcutaneous tissue, which leads to decreased adherence of the dermis to the fat in the early stages. 11

Unless an acute lesion is biopsied histology can be meaningless so that children presenting with a loss of subcutaneous fat are particularly difficult to diagnose. In the four patients described by Peters and Winkelmann nodules were not present at the beginning of the disease.
or if present were obvious only briefly.12 Three of their patients were thought to have lipo-
dystrophy until biopsies showed panniculitis.
Upper respiratory tract infections preceded
the panniculitis in two of our cases with lobular
panniculitis. The child with sepal panniculitis
suffered frequent exacerbations of her disease
after bouts of tonsillitis as did a child with
lobular panniculitis described by Gupta and
Rasmussen.13 Sepal panniculitis has been
recently associated with Lyme disease.14 There
is a similarity between this form of panniculitis
and erythema nodosum. Erythema nodosum is
considered to be a hypersensitivity reaction to a
variety of antigenic stimuli and is seen during
several infectious diseases including those
caused by β haemolytic streptococci. It is clear
from the literature that the presence of infection
is not predictive of the histological type of
panniculitis nor the clinical outcome.

The infant who underwent corrective cardiac
surgery was initially thought to have ‘cold
panniculitis’ resulting from the hypothermia
during surgery. A recent case report describes
an infant with similar problems after cardiac
surgery and the diagnosis of subcutaneous fat
necrosis of the newborn was made on the
histological findings.15 Cold panniculitis has
previously been described but is localised to the
area affected by the hypothermia, begins hours
to at most a few days after the insult, and does
not spontaneously recur without further
exposure to cold.16 Our child’s lesions recurred
too long after cardiac surgery to be consistent
with cold panniculitis. Nor is this child’s
condition compatible clinically or histologically
with sclerema neonatorum or subcutaneous fat
necrosis of the newborn.

Immunological abnormalities have previously
been reported in association with panniculitis. Allen-Mersh described a 7 year old girl
who first presented with a panniculitis of the foot
who went on to develop diabetes mellitus, hepatic cirrhosis, red cell surface antibodies,
and antibodies to mitochondria and gastric
parietal cells.17 Recently Billings et al
reported three cases of lobular panniculitis.18
One of the patients had insulin dependent
diabetes and Hashimoto’s thyroiditis, the
second later developed juvenile chronic arthritis,
and the third insulin dependent diabetes.

Hendricks et al describe a child whose disease
began after immunisation and another who did
not develop new lesions while he was lymphocytopen depl.

Our two older children with lobular panniculitis had positive rheumatoid factors. One had
a raised ASOT and immune complexes. All
children had negative autoimmune screen
The significance of these isolated findings is not clear and a review of the literature does not
suggest any correlation between the nature of
the immunological disturbance and the histo-
logical type of panniculitis.

It is clear that many factors are involved in
the initiation of panniculitis. It is possible that
various physical and chemical agents and
infections expose parts of the lipocyte increasing
its vulnerability to attack by autoimmune
mechanisms. This is supported by the increased
incidence of panniculitis among children with
t-antitrypsin deficiency.19

Steroids have been the mainstay of treatment
of panniculitis but some children do not
respond.20 Other agents tried with success
have been chloroquine;21 azathioprine,22
and cyclophosphamide.23

Panniculitis is a disease with at least two
distinct histological subtypes. Current treat-
ments are often based on the results presented
in numerous case reports rather than the
outcome of prospective trials of treatment. The
incidence of panniculitis is low but for a
particularly child this illness may cause serious
disability or even death. It is for these reasons
it is suggested that all children who present with
recurrent painful nodules of panniculitis have
an early and adequate biopsy to ascertain the
histology. Prompt treatment may prevent further
severe illness as well as disfiguring
subcutaneous fat loss.

1 Ackerman B. Panniculitis. Histologic diagnosis of inflammatory
2 Bagel J, Grossman ME. Subcutaneous nodules in pseudo-
3 Winkelmann RK. Panniculitis in connective tissue disease. Arch
4 Aronson JK, West DP, Vanakoski D, Ronan SG, Lonides I, Zeitz
HJ. Panniculitis associated with cutaneous T-cell lymphoma and
5 Edge J, Dunger DB, Dillon MJ. Weber-Christian panniculitis and
6 Hendricks WM, Ahmad M, Grazi E. Weber-Christian
7 Niemi KM, Forststrom L, Hannukela M, Mustakolto KK, Sala
OP. Nodules on the legs. A clinical, histological and
immunohistological study of 82 patients representing
different types of nodular panniculitis. Acta Derm Venereol
9 Taylor GA. Prolonged remission of Weber-Christian
10 Conway SP, Smithells RW, Peters WM, Weber-Christian
12 Peters MS, Winkelmann RK. Localised lipatrophy (atrophy
13 Gupta AK, Rasmussen JE. Multiple areas of localized tissue
14 Kramer N, Rickert RR, Brodkin RH, Rosenstein EO. Septal
panniculitis as a manifestation of Lyme disease. Am J Med
1986;81:149-52.
15 Silverman AK, Mickels EH, Rasmussen JE. Subcutaneous
fat necrosis in an infant occurring after hypothermic cardiac
16 Hultcrantz E. Harthuxen’s disease. Cold panniculitis in
17 Allen-Mersh TG. Weber-Christian panniculitis and auto-
18 Billings JK, Milgram SS, Gupta AK, Headington JT, Rasmussen
JE. Lipoatrophic panniculitis: a possible autoimmune inflammatory disease of fat. Arch
19 Rubenstein HM, Jaffer AM, Kudma JC, Lorrratanakul Y,
20 Winkelmann RK, McEvoy MT, Peters MS. Lipoplastic
21 Shelley WB. Chloroquine-induced remission of nodular
22 Kirch W, Duhren V, Hoenkel H, Ohnhaus E. Cyclo-
phosphamide-induced remission in Weber-Christian
23 Uplekar MW, Anisa NH. Dapsone dependent nodular
24 Roth DE, Schikler KS, Callen JP. Annular atrophic connective