The classification of paediatric vasculitis

Background to classification criteria

- Classification criteria are often described for diseases where the pathogenesis and/or molecular mechanisms are poorly understood.
- They are used to facilitate clinical trials and improve epidemiological descriptions by providing a set of agreed criteria that can be used by investigators anywhere in the world.
- Classification criteria for vasculitis are designed to differentiate one form of vasculitis from another once the diagnosis of vasculitis has been secured. They are not the same as diagnostic criteria (such as those described for Kawasaki disease), but are often misused as such.
- Thus, classification criteria aim to:
  - Identify a set of clinical findings (criteria) that recognize a high proportion of patients with the particular disease (sensitivity), and
  - Exclude a high proportion of patients with other diseases (specificity).
- Classification criteria typically include manifestations that are characteristics of the disease in question that occur with less frequency or are absent in other conditions.
- Symptoms or findings that might be typical or common but may also be present in other diseases tend to be excluded.
- An important limitation to these criteria is that they are not based on a robust understanding of the pathogenesis and as such are relatively crude tools that are likely to be modified as scientific understanding of these diseases progresses.

Paediatric vasculitis classification 2010

- New paediatric classification criteria are described, and validated on >1300 cases worldwide (Table 4.1).
- These criteria do not include Kawasaki disease (see Kawasaki disease, p. 183); nor do they include definitions for microscopic polyangiitis (too few cases included in dataset).
- For Takayasu arteritis, care must be taken to exclude fibromuscular dysplasia (or other cause of non-inflammatory large- and medium-vessel arteriopathy) since undoubtedly there could be scope for overlap in the clinical presentation between these 2 entities, although the pathogenesis and treatment for these are clearly distinct.

General scheme for the classification of paediatric vasculitides

- This is based on the size of the vessel predominantly involved in the vasculitic syndrome and is summarized as follows.
- It should be noted, however, that most vasculitides exhibit a significant degree of ‘polyangitis overlap’: e.g. Wegener’s granulomatosis can affect the aorta and its major branches, and small vessel vasculitis can occur in polyarteritis nodosa.
1. **Predominantly large-vessel vasculitis**
   - Takayasu arteritis.

2. **Predominantly medium-sized vessel vasculitis**
   - Childhood polyarteritis nodosa
   - Cutaneous polyarteritis
   - Kawasaki disease.

3. **Predominantly small-vessel vasculitis**
   - Granulomatous:
     - Wegener’s granulomatosis
     - Churg–Strauss syndrome
   - Non-granulomatous:
     - Microscopic polyangiitis
     - Henoch–Schönlein purpura
     - Isolated cutaneous leucocytoclastic vasculitis
     - Hypocomplementemic urticarial vasculitis.

4. **Other vasculitides**
   - Behçet’s disease
   - Vasculitis 2nd to infection (including hepatitis B-associated PAN), malignancies and drugs, including hypersensitivity vasculitis
   - Vasculitis associated with other connective tissue diseases
   - Isolated vasculitis of the CNS (childhood 1st angiitis of the central nervous system: cPACNS)
   - Cogan’s syndrome
   - Unclassified.
### Table 4.1 Classification criteria for specific vasculitic syndromes

<table>
<thead>
<tr>
<th>Vasculitis</th>
<th>Classification criteria</th>
<th>Sensitivity</th>
<th>Specificity</th>
</tr>
</thead>
<tbody>
<tr>
<td>HSP</td>
<td>Purpura, predominantly lower limb or diffuse(^a) (mandatory) plus 1 out of 4 of: Abdo pain, IgA on biopsy, Haematuria/proteinuria, Arthritis/arthralgia</td>
<td>100%</td>
<td>87%</td>
</tr>
<tr>
<td></td>
<td>(^a)If diffuse (i.e. atypical distribution) then IgA deposition on biopsy required</td>
<td></td>
<td></td>
</tr>
<tr>
<td>WG</td>
<td>At least 3 out of 6 of the following criteria: Histopathology, Upper airway involvement, Laryngo-tracheobronchial stenoses, Pulmonary involvement, ANCA positivity, Renal involvement</td>
<td>93%</td>
<td>99%</td>
</tr>
<tr>
<td>PAN</td>
<td>Histopathology or angiographic abnormalities (mandatory) plus 1 out of 5 of the following criteria: Skin involvement, Myalgia/muscle tenderness, Hypertension, Peripheral neuropathy, Renal involvement</td>
<td>89%</td>
<td>99%</td>
</tr>
<tr>
<td>TA</td>
<td>Angiographic abnormalities of the aorta or its main branches (also pulmonary arteries) showing aneurysm/dilatation (mandatory criterion), plus 1 out of 5 of the following criteria: Pulse deficit or claudication, 4 limb BP discrepancy, Bruits, Hypertension, Acute phase response</td>
<td>100%</td>
<td>99%</td>
</tr>
</tbody>
</table>


The epidemiology of paediatric vasculitis

- Childhood vasculitis is rare and the incidence and prevalence are not accurately described.
- Henoch–Schönlein purpura (HSP) and Kawasaki disease (KD) are the 2 commonest childhood vasculitides and as such those with the most epidemiological information.
  - There is undoubtedly some ethnic variation for these diseases (see individual section).
- In paediatric populations other systemic vasculitides including polyarteritis nodosa (PAN), Wegener’s granulomatosis (and other ANCA-associated vasculitides), Behçet’s disease, and Takayasu arteritis are rare and epidemiology difficult to assess.
  - Some ethnic variation has been noted: Takayasu’s being more common in Asians than North Americans and Europeans.

Henoch–Schönlein purpura
The estimated annual incidence in the West Midlands, UK has been reported as 20.4 per 100,000 and was highest in the 4–7yr age group. This is comparable to reported figures from the Czech Republic of 10.2 per 100,000 and Taiwan of 12.9 per 100,000.

Kawasaki disease
- KD has the highest incidence and prevalence in Asian populations, particularly Japan.
- Nationwide surveys conducted in Japan show the incidence of KD in children aged 0–4yr continues to rise with the average annual incidence in 2007 being 184.6 per 100,000.
  - This compares to an estimated annual incidence rate of 5.5–8.1 per 100,000 (0–5yr) in the UK and an estimated annual incidence rate of 1.6 per 100,000 (0–5yr) in the Czech Republic.
- There are limitations to these studies as they were all done by survey or questionnaire reporting.

Further reading
The investigation of primary systemic vasculitis

Background
Clinical features that suggest a vasculitic syndrome:
• Pyrexia of unknown origin
• Palpable purpura, urticaria, dermal necrosis
• Mononeuritis multiplex
• Unexplained arthritis, myositis, serositis
• Unexplained pulmonary, cardiovascular, or renal disease
• **Plus** 1 or more of:
  • Leucocytosis, eosinophilia
  • Hypocomplementaemia, cryoglobulinaemia
  • Circulating immune complexes
  • Raised ESR or CRP, thrombocytosis.

**Level 1 investigations—to be performed in all**
- Haematology and acute phase reactants:
  - FBC, ESR, CRP, clotting, prothrombotic screen (if patchy ischaemia of digits or skin), blood film
- Basic biochemistry:
  - Renal and liver function, CPK, thyroid function, LDH, amylase/lipase, urine dip and UA:UC (spot urine albumin:creatinine ratio)
- Infectious disease screen:
  - Blood cultures
  - Urine MC&S
  - ASOT and anti-DNase b
  - *Mycoplasma pneumoniae* serology
- Immunological tests:
  - ANA, dsDNA Abs, ENAs, ANCA, RF,
  - Anti-GBM antibodies
  - TTG antibodies (coeliac disease screen)
  - Immunoglobulins: IgG, IgA, IgM, and IgE
  - Anticardiolipin antibodies, lupus anticoagulant
  - C3/C4, MBL (memose binding lectin, if available), CH100 or alternative functional complement assay if available
  - VZV antibody status (prior to starting immunosuppressive therapy)
  - Serum ACE
- Radiological: CXR, abdominal and renal USS
- Other: ECG, echocardiography, digital clinical photography of lesions.

**Level 2 investigations—to be considered on an individual basis**
- Infection screen:
  - Mantoux 1:1000, and/or quantiferon
  - PCR for CMV, EBV, enterovirus, adenovirus, VZV, HBV, HCV
  - Serology for HIV, Rickettsiae, *Borrelia burgdorferi*
• Imaging:
  • Radiograph of bones and joints.
  • Selective contrast visceral angiography.
  • DMSA scan.
  • MRI/MRA of brain (for suspected cerebral vasculitis).
  • CT abdomen, thorax, brain, sinus X ray (for Wegener’s).
  • Labelled white cell scan (for extent and location of inflammation).
  • Cerebral contrast angiography (for suspected cerebral vasculitis).
  • PET-CT: for differential of malignancy or Castleman’s disease.
  • DEXA scan.
  • V/Q scan.
  • USS Doppler of peripheral arteries.
  • Thermography and nail fold capillaroscopy.
• Tissue biopsy: skin, nasal or sinus, kidney, sural nerve, lung, liver, gut, temporal artery, brain, other.
• Bone marrow analysis and/or lymph node excision biopsy (for suspected malignancy).
• Biochemistry, immunology and immunogenetics:
  • Serum amyloid A.
  • Formal GFR.
  • Organ specific autoantibodies.
  • IgD.
  • B2 Glycoprotein 1 antibodies.
  • Urinary catecholamines (consider plasma catecholamines as well), and urine VMA, HVA (for phaeochromocytoma, or neuroblastoma).
  • Cryoglobulins (if there is a history of cold sensitivity/vasculitis mainly present in exposed areas of the body).
  • Basic lymphocyte panel and CD19 count if monitoring post rituximab.
  • Mitochondrial DNA mutations.
  • DNA analysis for periodic fever syndromes that can mimic vasculitis: MEFV (familial Mediterranean fever, TNFRSF1A (TNFα receptor associated periodic fever syndrome, TRAPS), MVK (hyper IgD syndrome, HIDS), NLRP3 (cryopyrin associated periodic syndrome, CAPS), and NOD2 (Crohn’s/Blau’s/juvenile sarcoid mutations).
  • Nitroblue tetrazolium test if granulomatous inflammation found on biopsy.
• Nerve conduction studies (PAN, WG, Behçet’s [before starting thalidomide]).
• Ophthalmology screen.
• Ambulatory 24h BP/4-limb BP.
The standard treatment of childhood vasculitis

Guidelines for the use and monitoring of cytotoxic drugs in non-malignant disease are shown in Table 4.2. Standard vasculitis therapy (excluding crescentic glomerulonephritis) is described in Fig. 4.1. Prior to using this approach, remember there should be:

- A well-established diagnosis.
- Severe, potentially life-threatening disease.
- Inadequate response to less toxic therapy—milder cases of vasculitis (e.g. isolated cutaneous forms) may respond to less toxic agents such as colchicine. Therapy should always be tailored for each individual.
- No known infection or neoplasm.
- No pregnancy or possibility thereof.
- Informed consent obtained and documented in notes.

Other points of note

- Although the use of oral cyclophosphamide is highlighted in Table 4.3, increasingly IV cyclophosphamide is favoured over the oral route in children and adults because of reduced side effects and lower cumulative dose, but comparable efficacy as suggested by a number of studies in adults with ANCA vasculitis (e.g. the 'CYCLOPS' trial).
- IV cyclophosphamide has the added advantage of ensuring adherence to therapy, of particular relevance in adolescents with vasculitis.

Use of biologic therapy in systemic vasculitis of the young

- Whilst the therapeutic approach and drugs used as suggested in Figs. 4.1 and 4.2 undoubtedly have improved survival and long-term outlook for children with severe vasculitis, concerns relating to toxicity particularly with cyclophosphamide, and relapses despite this conventional therapeutic approach have led to the increasing use of biologic therapy such as rituximab, anti-TNF alpha, or other biologic therapy.
- Evidence to support the use of rituximab as a 1st induction agent in place of cyclophosphamide for the treatment of ANCA-associated vasculitis is now available for adults with this group of diseases (RITUXIVAS, and RAVE trials).
- Evidence to support this approach in children remains anecdotal, but undoubtedly rituximab is being increasingly used for children with ANCA vasculitis that is not adequately controlled using the conventional cyclophosphamide followed by azathioprine therapeutic regimen outlined in Fig. 4.1.
- Evidence for the use of anti-TNFα or other biologic agents such as anakinra remains anecdotal for children and adults with vasculitis.
- Whilst there is not enough evidence to recommend specific biologic therapy for specific vasculitic syndromes, a general approach favoured by the author is given in Table 4.3.
Table 4.2 Doses, side effects, and clinical monitoring of commonly used immunosuppressant and cytotoxic immunosuppressant drugs used for the treatment of vasculitis

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dose</th>
<th>Side effects</th>
<th>Cumulative toxic dose</th>
<th>Clinical monitoring</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cyclophosphamide (CYC)</td>
<td>2–3mg/kg once a day PO 2–3 months; 0.5–1.0g/m² IV monthly with mesna to prevent cystitis (see Chapter 9 for mesna dose and IV CYC administration protocol)</td>
<td>Leucopenia; haemorrhagic cystitis; reversible alopecia; infertility; leukaemia, lymphoma, transitional cell carcinoma of bladder</td>
<td>Not described for malignancy; 500mg/kg for azoospermia</td>
<td>Weekly FBC for duration of therapy (usually 2–3 months); baseline and monthly renal and liver function. Temporarily discontinue and/or reduce dose if neutropenia &lt;1.5 × 10⁹/L, platelets &lt;150 × 10⁹/L, or haematuria. Day 10 FBC if IV. Reduce dose if renal or hepatic failure e.g. to 250–300mg/m².</td>
</tr>
<tr>
<td>Azathioprine (600 mg/m² twice a day)</td>
<td>0.5–2.5mg/kg once a day PO for 1yr or more</td>
<td>GI toxicity; hepatotoxicity; rash; leucopenia; teratogenicity; no increase in malignancy in adults with RA; no conclusive data for cancer risk in children</td>
<td>Not described</td>
<td>Weekly FBC for 1 month, then 3-monthly. Temporarily discontinue and/or reduce dose if neutropenia &lt;1.5 × 10⁹/L, platelets &lt;150 × 10⁹/L, and check TPMT enzyme—patients deficient in TPMT require reduced doses (or may not tolerate) azathioprine because of marrow toxicity.</td>
</tr>
<tr>
<td>Mycophenolate mofetil (MMF)</td>
<td>(600 mg/m² twice a day)</td>
<td>Bone marrow suppression; severe diarrhoea; pulmonary fibrosis</td>
<td>Not described</td>
<td>Fortnightly FBC for 2 months, then monthly for 2/12. 3-monthly when stable. Baseline monthly renal and liver function until stable. Discontinue temporarily and/or reduce dose if neutropenia &lt;1.5 × 10⁹/L, platelets &lt;150 × 10⁹/L, or significant GI side effects.</td>
</tr>
<tr>
<td>Ciclosporin</td>
<td>3–5mg/kg/day PO in 2 divided doses</td>
<td>Renal impairment, hypertension, hepatotoxicity, tremor, gingival hyperplasia, hypertrichosis, lymphoma</td>
<td>Not described</td>
<td>Weekly measurement of BP; baseline then monthly renal and liver function; maintain 12h trough level at 50–100ng/mL. 6–12-monthly GFR. Consider renal biopsy every 2 years.</td>
</tr>
<tr>
<td>Methotrexate</td>
<td>10–15mg/m²/week PO or SC (single dose)</td>
<td>Bone marrow suppression and interstitial pneumonitis (risk with folic acid), reversible elevation of transaminases, hepatic fibrosis</td>
<td>Not described</td>
<td>Baseline CXR, FBC, and LFTs, then FBC and LFTs fort nightly until dose stable, then monthly to every 6 weeks (after 6 months). Reduce or discontinue if hepatic enzymes &gt;3× upper limit of normal, neutropenia &lt;1.5 × 10⁹/L, new or worsening cough, severe nausea, vomiting, or diarrhoea, platelets &lt;150 × 10⁹/L or falling rapidly.</td>
</tr>
</tbody>
</table>
Induction therapy
• Prednisolone 30–60 mg/m² once a day (1–2 mg/kg) for 4 weeks, weaning over next 6–8 weeks (depending on response to Rx) to 0.3–0.7 mg/kg on alternate days; or IV methylprednisolone 30 mg/kg (max. 1g) for 3 consecutive days followed by oral prednisolone as above.
• CYC 2–3 mg/kg PO once a day for 2–3/12 or 500–1000 mg/m² IV (max. 1.2 g) once a month for 6 months (reduce dose if renal or hepatic failure). Aspirin 1–2 mg/kg once a day (or dipyridamole 2.5 mg/kg BD if aspirin contraindicated).

Failed induction

Post induction (maintenance) phase: 18 months to 3 yr for PAN; may require prolonged Rx in some vasculitic syndromes.
• Azathioprine 2–3 mg/kg PO once a day (start 3–5 days after stopping PO CYC; 10 days after IV CYC); consider measuring TPMT first.
• Prednisolone 0.2–0.5 mg/kg alternate days (daily if ongoing disease activity).
• Aspirin 1–2 mg/kg once a day or dipyridamole 2.5 mg/kg twice a day
• Consider ranitidine or proton pump inhibitor.

Consider
• Single dose of IV CYC 750–1000 mg/m² if previously given oral CYC for induction of remission.
• Methylprednisolone 30 mg/kg (max. 1g) IV x3
• 5- or 10-day course of daily 2-volume plasma exchange with 4.5% HAS.
• 2nd course of oral CYC 2 mg/kg once a day for 2 weeks.
• IVIG 2 g/kg.
• Biologic agent:
  • Anti-TNF therapy
  • Rituximab

Major relapse whilst on maintenance therapy.

Notes:
1. 2nd-line maintenance agents
   1. MMF
   2. Ciclosporin
   3. MTX
   4. Colchicine
2. Consider sperm cryopreservation for all post-pubertal males receiving CYC.
3. For monitoring of complications of therapy refer to Table 4.2.
4. Beware neutropenia as prednisolone dose is weaned during maintenance phase of therapy.
5. Miscellaneous vasculitides such as Behçet’s may require colchicine and/or thalidomide.
6. Treatment with biological agents in select individuals who fail to respond to standard induction therapy (see separate guidelines).
7. Epoprostenol (prostacyclin) 1–20 ng/kg/min IV for incipient gangrene
8. Other agents with as yet unproven efficacy in childhood vasculitis: leflunamide; DSG

Minor relapse: ↑ oral prednisolone
• Recurrent minor relapses or ‘grumbling vasculitis’: consider IV pulsed methylprednisolone and/or switch to 2nd-line maintenance therapy.

Stopping treatment
• Usually withdrawn slowly over 6 months if no relapse for 12 months.
• Recommend stopping azathioprine first over 3 months, followed by gradual taper of prednisolone over next 3 months.

Fig. 4.1 Standard vasculitis therapy (excluding crescentic glomerulonephritis).
Fig. 4.2 Guidelines for treatment of crescentic glomerulonephritis (note early diagnosis and starting therapy is of major importance).

- **Linear IgG staining on immunofluorescence**
  - Anti GBM +
  - Treatment as per vasculitis algorithm above, but consider using plasma exchange as first-line induction therapy in conjunction with steroid and CYC

- **Pauci-immune on immunofluorescence**
  - Consider microscopic polyangiitis or ‘renal-limited vasculitis’—may be ANCA +

- **Granular deposits on immunofluorescence**
  - Indicative of immune complex disease
  - Electron microscopy

- **Crescentic GN on renal biopsy**
  - Plasma exchange for 10–14 days (2 volume, 4.5% HAS)—or until anti GBM Ab disappears
  - Pulsed IV methyl prednisolone 30 mg/kg (max. 1 g) x3, then oral prednisolone 2 mg/kg once a day (weaned over 2 months then stop)
  - CYC 2 mg/kg PO once a day for 2 months, or IV at 500 mg/m² monthly (reduced dose because of renal failure), possibly increasing by 250 mg/m² per month (response dependent) to maximum of 1000 mg/m² (max. 1.2 g) for 6 months
  - Consider prolonging therapy if Anti GBM still detectable

- **Failed remission**
  - Consideration of biologic therapy:
    - Rituximab
    - Anti-TNF therapy
  - Refer to specific protocols

- **Consider treating specific disorders (refer to relevant protocols):**
  - HSP
  - IgA nephropathy
  - Post streptococcal
  - SLE
  - Membranoproliferative GN
  - Membranous GN
### Table 4.3

Recommendations for indication and choice of biologic therapy for primary systemic vasculitis of the young based on published experience

<table>
<thead>
<tr>
<th>Vasculitis type</th>
<th>Indication for biologic agent</th>
<th>Proposed first choice of biologic agent</th>
</tr>
</thead>
<tbody>
<tr>
<td>ANCA-associated vasculitis</td>
<td>Critical organ or life-threatening disease which has failed to respond to standard vasculitis therapy or concerns regarding cumulative CYC dose.</td>
<td>Rituximab or other B cell depleting monoclonal antibody</td>
</tr>
<tr>
<td>Polyarteritis nodosa</td>
<td>Failed therapy with standard agents or concern regarding cumulative CYC dose</td>
<td>Rituximab or anti-TNFα†</td>
</tr>
<tr>
<td>Behçet’s disease</td>
<td>Recalcitrant and severe disease; alternative to thalidomide</td>
<td>Anti-TNFα (infliximab, adalimumab or etanercept)</td>
</tr>
</tbody>
</table>

CYC, cyclophosphamide; †Authors have more experience with infliximab than etanercept for PAN, although etanercept has been used in individual cases of childhood PAN; ††No firm recommendation is made regarding first choice of biologic for PAN.


### Further reading

Henoch–Schönlein purpura

Introduction
- Henoch-Schönlein purpura (HSP), the commonest systemic vasculitis in childhood is predominantly a small-vessel, non-granulomatous leucocytoclastic vasculitis of unknown aetiology.
  - Reported by Heberden (1801), Willan (1808), Schönlein (1832), and Henoch (1874).
  - Also called anaphylactoid purpura (1948).
- Classification criteria are palpable purpura in a predominantly lower limb distribution with at least 1 of 4 of:
  - Diffuse abdominal pain.
  - Any biopsy showing IgA deposition (mandatory criterion if rash is atypical).
  - Arthritis and/or arthralgia.
  - Haematuria and/or proteinuria.
- Variable and often relapsing course without specific laboratory findings with 1/3 of children having symptoms up to a fortnight, another 1/3 up to 1 month and recurrence of symptoms within 4 months of resolution in 1/3.
- Henoch–Schönlein nephritis (HSN) accounts for 1.6–3% of all childhood cases of end-stage renal failure (ESRF) in the UK.

Epidemiology
- Commoner in Caucasian and Asian populations and boys: 
  - O:O ratio of 1.5–2:1.
  - 50% present before the age of 5yr, 75% present before the age of 10yr.
- Incidence of 10–20.4 (mean of 13.5) per 100,000 children.
  - 22.1 per 100,000 children under 14yr of age.
  - 70.3 per 100,000 children aged 4–7yr.
- Almost 20× rarer in adults.
  - 0.8 cases per 100,000 adults.
  - More severe in adults.
- Seasonal variation (commoner in winter) with infectious triggers.
  - Associations with bacteria (e.g. Group A beta-haemolytic streptococci) and viruses (hepatitis, CMV, HSV, human parvovirus B19, coxsackie, and adenovirus), and some cases after vaccination described.

Immunopathology
- Type III hypersensitivity reaction with immune complex.
  - IgA deposition: galactose deficient IgA may contribute to this.
  - Alternate pathway complement activation.
  - Associated C2 deficiency.
- Vasculitis of small blood vessels with diffuse angiitis.
  - Perivascular exudate of leucocytes.
- Polygenic inheritance with renal involvement associated with HLA-B35, IL-1β (−511) T allele, and IL-8 allele A.
CHAPTER 4 Systemic diseases

Systemic involvement

Dermatological
- All patients have purpura but skin involvement may not be present at time of initial presentation.
- Generally symmetrical purpura involving lower limbs and buttocks but can spread to upper limbs (rarely abdomen, chest or face).
- Urticaria and angio-oedema can occur.

Gastrointestinal
- Commonly occurs (68% of patients).
- Abdominal pain precedes rash by 1–14 days in 43% of patients.
- Presents with intermittent colicky abdominal pain, vomiting with or without haematemesis or melaena (faecal occult blood may be positive) as a result of haemorrhage into gut wall.
- Involvement may result in intussusception, appendicitis, cholecystitis, pancreatitis, GI haemorrhage, ulceration, infarction, or perforation.

Rheumatological
60% of patients will have joint involvement with arthritis and/or arthralgia usually affecting the knees and ankles resulting in pain, swelling and ↓ range of movement.

Renal
- 25–60% of patients will have renal involvement with HSN:
  - 76% will develop within 4 weeks of disease onset.
  - 97% will develop within 3 months of disease onset.
- Most cases are usually asymptomatic which necessitates screening up to 6 months after last recrudescence of rash or HSP symptoms with <10% having significant involvement requiring referral for consideration of renal biopsy (Fig. 4.3).
  - Microscopic haematuria without proteinuria is benign.
  - 82% have normal renal function after 23yr.
  - Good prognosis with isolated haematuria and mild proteinuria with mild histological changes as less than 5% will develop chronic kidney disease (CKD) within 10–25yr.
  - Improved renal prognosis in children <7yr.
- Severe disease indicated by increasing proteinuria, development of nephrotic syndrome and/or renal failure.
  - 20% of patients with acute mixed nephritic and nephrotic syndrome progress to ESRF.
  - 44–50% develop hypertension or CKD.
  - Histological pattern is identical to IgA nephropathy and includes focal segmental proliferative glomerulonephritis and rapidly progressive crescentic glomerulonephritis (Table 4.4).

Other
Patients may present with orchitis, severe pulmonary haemorrhage, and/or cerebral vasculitis (which may respond to immunosuppression combined with plasmapheresis).
Treatment
- Symptomatic treatment with rest and analgesia.
- No role of antibiotics unless suspected or proven infection.
- Prophylactic corticosteroid therapy at commencement of HSP does not prevent renal or GI involvement.
- However, corticosteroids do seem to be effective in treating these complications and severe facial and/or scrotal haemorrhagic oedema.
- Patients with severe renal involvement may require other immunosuppressive agents, antiproteinuric and antihypertensive agents.

<table>
<thead>
<tr>
<th>ISKDC grade</th>
<th>Pathoanatomical findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Minimal alterations</td>
</tr>
<tr>
<td>II</td>
<td>Mesangial proliferation</td>
</tr>
<tr>
<td>III A</td>
<td>Focal proliferation or sclerosis with &lt;50% crescents</td>
</tr>
<tr>
<td>III B</td>
<td>Diffuse proliferation or sclerosis with &lt;50% crescents</td>
</tr>
<tr>
<td>IV A</td>
<td>Focal proliferation or sclerosis with 50–75% crescents</td>
</tr>
<tr>
<td>IV B</td>
<td>Diffuse proliferation or sclerosis with 50–75% crescents</td>
</tr>
<tr>
<td>V A</td>
<td>Focal proliferation or sclerosis with &gt;75% crescents</td>
</tr>
<tr>
<td>V B</td>
<td>Diffuse proliferation or sclerosis with &gt;75% crescents</td>
</tr>
<tr>
<td>VI</td>
<td>Membranoproliferative glomerulonephritis</td>
</tr>
</tbody>
</table>

ISKDC=International Study of Kidney Diseases in Children.
Fig. 4.3 Clinical pathway for investigation and referral for renal biopsy in HSN: Reproduced with permission from Tizard EJ, Hamilton-Ayres MJ.
Henoch–Schönlein purpura. Arch Dis Child Educ Pract Ed 2008; 93:1–8.AIP, auto-immune profile; ANCA, anti-neutrophil cytoplasmic Abs; ARF, acute renal failure; ASOT, anti-streptolysin O titre; BP, blood pressure; C3,C4, complement C3 and C4; EMU, early morning urine; GP, general practitioner; Ht, height; U&E, urea and electrolytes; Wt, weight; UP:UC, urine protein:creatinine ratio.

Indications for consideration of renal biopsy:
1. Acute nephritic syndrome/ARF
2. Nephrotic syndrome/nephrotic range proteinuria (UP:UC>250 mg/mmol) for 4–6 weeks

* Review in primary care/with local paediatrician according to local arrangements NB children with recurrent episodes should monitored as for the first episode.

W1–W4: weekly GP* review (BP, EMU dipstick)

WS–W12: fortnightly GP* review (BP, EMU dipstick)

6–12: month GP* review (BP, EMU dipstick)
Kawasaki disease

Kawasaki disease (KD) is a self-limiting vasculitic syndrome that predominantly affects medium- and small-sized arteries. It is the 2nd commonest vasculitic illness of childhood (the commonest being HSP) and it is the leading cause of childhood acquired heart disease in developed countries.

Pathogenesis and epidemiology

- Pronounced seasonality and clustering of KD cases have led to the hunt for infectious agents as a cause. However, so far no single agent has been identified.
- The aetiology of KD remains unknown but it is currently felt that one or more widely distributed infectious agents evoke an abnormal immunological response in genetically susceptible individuals, leading to the characteristic clinical presentation of the disease.
- KD has a world-wide distribution with a 4 preponderance, an ethnic bias towards Asian and in particular Japanese or Chinese children, some seasonality, and occasional epidemics.
- The incidence of KD is rising world-wide, including the UK. The current reported incidence in the UK is 8.1/100,000 children aged <5yr. This may reflect a truly rising incidence or † clinician awareness.

Clinical features

The principal clinical features of KD are:
- Fever persisting for 5 days or more.
- Peripheral extremity changes (reddening of the palms and soles, indurative oedema, and subsequent desquamation).
- A polymorphous exanthema.
- Bilateral conjunctival injection/congestion.
- Lips and oral cavity changes (reddening/cracking of lips, strawberry tongue, oral and pharyngeal injection), and
- Acute non-purulent cervical lymphadenopathy.

- For the diagnosis of KD to be established 5 of the 6 clinical features should be present.
- Patients with <5 or 6 principal features can be diagnosed with KD when coronary aneurysm or dilatation is recognized by two-dimensional (2D) echocardiography or coronary angiography.
- The cardiovascular features are the most important manifestations of the condition with widespread vasculitis affecting predominantly medium-size muscular arteries, especially the coronary arteries. Coronary artery involvement occurs in 15–25% of untreated cases with additional cardiac features in a significant proportion of these including pericardial effusion, electrocardiographic abnormalities, pericarditis, myocarditis, valvular incompetence, cardiac failure, and myocardial infarction.
- Of note, irritability is an important sign, which is virtually universally present although not included in the diagnostic criteria.
- Another clinical sign that may be relatively specific to KD is the development of erythema and induration at sites of BCG inoculations. The mechanism of this sign is thought to be cross reactivity of T cells
in KD patients between specific epitopes of mycobacterial and human heat shock proteins.

- An important point worthy of emphasis is that the principal symptoms and signs may present sequentially such that the full set of criteria may not be present at any one time. Awareness of other non-principal signs (such as BCG scar reactivation) may improve the diagnostic pick-up rate of KD.
- Other clinical features include: arthritis, aseptic meningitis, pneumonitis, uveitis, gastroenteritis, meatitis and dysuria, and otitis.
- Relatively uncommon abnormalities include hydrops of the gallbladder, GI ischaemia, jaundice, petechial rash, febrile convulsions, and encephalopathy or ataxia, macrophage activation syndrome, and syndrome of inappropriate anti-diuretic hormone secretion (SIADH).

**Differential diagnosis**

Conditions that can cause similar symptoms to KD and must be considered in the differential diagnosis include:

- Scarlet fever
- Rheumatic fever
- Streptococcal or staphylococcal toxic shock syndrome
- Staphylococcal scalded skin syndrome
- Systemic JIA
- Infantile PAN
- SLE
- Adenovirus, enterovirus, Epstein–Barr virus, CMV, parvovirus, influenza virus infection
- *Mycoplasma pneumoniae* infection
- Measles
- Leptospirosis
- Rickettsiae infection
- Adverse drug reaction
- Mercury toxicity (acrodynia)
- Lymphoma—particularly for IVIg resistant cases.

**Investigations**

In cases of suspected KD the following investigations should be considered:

- FBC and blood film.
- ESR.
- CRP.
- Blood cultures.
- ASOT and anti-DNase B.
- Nose and throat swab, and stool sample for culture (superantigen toxin typing if *Staphylococcus aureus* and/or beta-haemolytic streptococci detected).
- Renal and liver function tests.
- Coagulation screen.
- Autoantibody profile (ANA, ENA, RF, ANCA).
- Serology (IgG and IgM) for *Mycoplasma pneumoniae*, enterovirus, adenovirus, measles, parvovirus, Epstein–Barr virus, cytomegalovirus.
- Urine MC&S.
Dip test of urine for blood and protein.
Consider serology for rickettsiae and leptospirosis if history suggestive.
Consider CXR.
ECG.
2D echocardiography to identify coronary artery involvement acutely and monitoring changes long term.
Coronary arteriography has an important role for delineating detailed anatomical injury, particularly for children with giant coronary artery aneurysms (>8mm), where stenoses adjacent to the inlet/outlet of the aneurysms are a concern. Note that the procedure may need to be delayed until at least 6 months after disease onset since there could be a risk of myocardial infarction if performed in children with ongoing severe coronary artery inflammation.

**Treatment (Fig. 4.4)**

The treatment of KD comprises of:
- IV Ig at a dose of 2g/kg as a single infusion over 12h (consider splitting the dose over 2–4 days in infants with cardiac failure).
- IV Ig should be started early preferably within the first 10 days of the illness. However, clinicians should not hesitate to give IV Ig to patients who present after 10 days if there are signs of persisting inflammation.
- Aspirin 30–50mg/kg/day in 4 divided doses.
- The dose of aspirin can be reduced to 2–5mg/kg/day when the fever settles (disease defervescence). Aspirin at antiplatelet doses is continued for a minimum of 6 weeks.
- If the symptoms persist within 48h or there is disease recrudescence within 2 weeks a 2nd dose of IV Ig at 2g/kg over 12h should be considered.
- However, IV Ig resistance occurs up to 20% of cases.
- When a patient fails to respond to a 2nd dose of IV Ig, consider IV pulsed methylprednisolone at 15–30mg/kg daily for 3 days to be followed by oral prednisolone 2mg/kg/day once a day weaning over 6 weeks. Some clinicians are increasingly using corticosteroids after disease recrudescence following one dose of IV Ig based on the results of a recent study. This remains an area of controversy, but seems rational since this is associated in most cases with rapid resolution of inflammation.
- In refractory cases infliximab, a human chimeric anti-TNFα monoclonal antibody, given IV at a single dose of 6mg/kg has been reported to be effective, and is increasingly used for IV Ig resistant cases. Considering that rapid and effective interruption of inflammation is a 1st target of KD therapy, TNFα blockade may be a logical step following one failed dose of IV Ig, particularly in very active disease with evidence of early coronary artery dilatation.
- Echocardiography should be repeated at 2 weeks and 6 weeks from initiation of treatment (refer to paediatric cardiology).
- If the repeat echocardiogram shows no coronary artery abnormalities (CAAs) at 6 weeks, aspirin can be discontinued and lifelong follow-up at least every 2yr should be considered.
In cases of CAA <8mm with no stenoses present, aspirin should be continued until aneurysms resolve.

If CAA >8mm and/or stenoses is present, aspirin at a dose of 2–5mg/kg/day should be continued lifelong. The combination of aspirin and warfarin therapy in these patients with giant aneurysms has been shown to ↓ the risk of myocardial infarction.

In patients who develop CAA, echocardiography and ECG should be repeated at 6-monthly intervals and an exercise stress test considered.

Other specific interventions such as PET scanning, addition of calcium channel blocker therapy, and coronary angioplasty should be organized at the discretion of the paediatric cardiologist.

**Outcome**

- Treatment with IVIg and aspirin reduces CAA from 25% for untreated cases to 4–9%.
- IVIg resistance occurs in approximately 20%, and is associated with a higher risk of CAA.
- The overall outlook of children with KD is good, with the acute mortality rate due to myocardial infarction having been reduced to <1% by ↑ alertness of the clinicians to the diagnosis and prompt treatment.
- Nonetheless the disease may contribute to the burden of adult cardiovascular disease and cause premature atherosclerosis, an area of active ongoing research.
**Fig. 4.4** Guideline for the management of Kawasaki disease.

1. **Establish diagnosis of Kawasaki disease**
   - IVIG 2g/kg as a single infusion over 12 h (consider splitting the dose over 2–4 days in infants with cardiac failure)
   - Aspirin 30–50 mg/kg/day in 4 divided doses
   - Perform echocardiography, and ECG
   - Aspirin 2–5 mg/kg/day when fever settled (disease defervescence) continuing for a minimum of 6 weeks

2. **Disease defervescence**
   - Repeat echocardiography at 2 weeks and 6 weeks

3. **No disease defervescence within 48 h, or disease recrudescence within 2 weeks**
   - a) 2nd dose of IVIG at 2g/kg over 12 h
   - Pulsed IV methyl prednisolone at 15–30 mg/kg daily for 3 days to be followed by oral prednisolone 2 mg/kg/day once a day weaning over 6 weeks—seek expert advice
   - Infliximab (6 mg/kg) for refractory cases—seek expert advice

4. **No coronary artery abnormalities (CAAs)**
   - Stop aspirin at 6 weeks
   - Consider lifelong follow-up at least every 2 years
   - CAA <8 mm, no stenoses
     - Continue aspirin until aneurysms resolve
     - Repeat echocardiography & ECG at 6-monthly intervals
     - Discontinue aspirin if aneurysms resolve
     - Consider exercise stress test if multiple aneurysms
     - Specific advice re: minimizing atheroma risk factors, and consider lifelong follow-up
   - CAA >8 mm, and/or stenoses
     - Lifelong aspirin 2–5 mg/kg/day
     - Warfarin (with initial full heparinization to prevent paradoxical thrombosis)
     - Consider coronary angiography (after at least 6 months from disease onset) and exercise stress testing
     - Repeat echocardiography & ECG at 6-monthly intervals
     - Specific advice re: minimizing atheroma risk factors
     - Lifelong follow-up

5. **Seek expert advice to consider**
   - Discontinue aspirin if aneurysms resolve
   - Consider exercise stress test if multiple aneurysms
   - Specific advice re: minimizing atheroma risk factors
   - Lifelong follow-up

**Notes:**

- Treatment can be commenced before full 5 days of fever if sepsis excluded; treatment should also be given if the presentation is >10 days from fever onset
- Refer to paediatric cardiologist
- Other specific interventions such as PET scanning, addition of calcium channel blocker therapy, and coronary angioplasty at discretion of paediatric cardiologist.
The anti-neutrophil cytoplasmic antibody (ANCA)-associated vasculitides

Background
• The ANCA-associated vasculitides (AAV) are:
  • Wegener’s granulomatosis (WG)
  • Microscopic polyangiitis (MPA)
  • Churg–Strauss syndrome (CSS) and
  • Renal limited vasculitis (previously referred to as idiopathic crescentic glomerulonephritis).
• Although rare, the AAV do occur in childhood.

Definitions of AAV
Definitions for each of the AAV describing the salient major clinical and laboratory features are given here. These are not the same as classification criteria, which (for WG) are provided in a separate section on classification.
• WG: granulomatous inflammation involving the respiratory tract and necrotizing vasculitis affecting small- to medium-size vessels
• MPA: necrotizing vasculitis, with few or no immune deposits, affecting small vessels; necrotizing arteritis involving small- and medium-sized arteries may be present; pulmonary capillaritis often occurs. Clinically, it often presents with rapidly progressive pauci-immune glomerulonephritis, in association with perinuclear ANCA (pANCA, MPO-ANCA) positivity.
• CSS: an eosinophil-rich and granulomatous inflammation involving the respiratory tract and necrotizing vasculitis affecting small- to medium-sized vessels; there is an association with asthma and eosinophilia.
• Renal limited: rapidly progressive glomerulonephritis, often with ANCA positivity (usually MPO-ANCA) but without other organ involvement.

Pathogenesis
• It is not known why patients develop ANCA in the first instance.
• When ANCA are present, the most accepted current model of pathogenesis proposes that ANCA activate cytokine-primed neutrophils, leading to bystander damage of endothelial cells and an escalation of inflammation with recruitment of mononuclear cells.
• However, other concomitant exogenous factors and genetic susceptibility appear to be necessary for disease expression.

Clinical features of WG
From a clinical perspective WG may be broadly considered as having 2 forms:
• Predominantly granulomatous form with mainly localized disease, and
• Florid, acute small vessel vasculitic form characterized by severe pulmonary haemorrhage and/or rapidly progressive vasculitis or other severe vasculitic manifestation.
These 2 broad presentations may coexist or present sequentially in individual patients.
Organ specific involvement includes:

- **Upper respiratory tract:**
  - Epistaxis.
  - Otalgia, and hearing loss (conductive and/or sensorineural); chronic otitis media; mastoiditis.
  - Nasal septal involvement with cartilaginous collapse results in the characteristic saddle nose deformity (Fig. 4.5).
  - Chronic sinusitis.
  - Glottic and subglottic polyps and/or large- and medium-sized airway stenosis.

- **Lower respiratory tract manifestations include (singly or in combination):**
  - Granulomatous pulmonary nodules with or without central cavitation.
  - Pulmonary haemorrhage with respiratory distress, frank haemoptysis, and/or evanescent pulmonary shadows (CXR).

- **Renal involvement:** typically a focal segmental necrotizing glomerulonephritis, with pauci-immune crescentic glomerular changes. The clinical manifestations associated with this lesion are:
  - Hypertension.
  - Significant proteinuria.
  - Nephritic and nephrotic syndrome.
  - Other protean manifestations of renal failure.

- **Ophthalmological disease:** retinal vasculitis, conjunctivitis, episcleritis, uveitis, optic neuritis. Unilateral or bilateral proptosis may be caused by granulomatous inflammation affecting the orbit (pseudotumour) (Fig. 4.5).

- **Malaise, fever, weight loss or growth failure, arthralgia, and arthritis.**

- **Other manifestations include:** peripheral gangrene with tissue loss, and vasculitis of the skin, gut (including appendicitis), heart, central nervous system and/or peripheral nerves (mononeuritis multiplex), salivary glands, gonads, and breast.

**Investigations (also see [Vasculitis investigation, p 172])**

- **WG** is commonly associated with a cytoplasmic staining pattern of ANCA by IIF, and ELISA reveals specificity against PR3 (PR3-ANCA).
- **MPA** and renal limited AAV are typically associated with pANCA by IIF and with MPO-ANCA specificity on ELISA.
- **ANCA-negative forms of WG, MPA, renal limited vasculitis, and CSS** are well described in children.
- **While the diagnostic value of ANCA is without question important,** the value of ANCA for the longitudinal monitoring of disease activity is probably unreliable in many patients with WG.
- **Tissue diagnosis,** in particular renal biopsy but also biopsy of skin, nasal septum, or other tissue, can be important diagnostically for diagnosing all of the AAV and can help stage the disease for therapeutic decision-making.
Other commonly observed non-specific findings include:
- Mild normochromic normocytic anaemia together with a leucocytosis and thrombocytosis.
- Elevated ESR and CRP.
- Raised immunoglobulins (polyclonal IgG).

Laboratory manifestations relating to renal involvement include:
- Dipstick haematuria and proteinuria positive.
- Raised urinary spot protein creatinine ratio.
- Raised serum creatinine and other associated laboratory features of renal failure.

Chest radiography may be abnormal but high resolution CT chest has better sensitivity for demonstrating pulmonary infiltrates or discrete nodular and/or cavitating lesions.

Plain x-ray or CT sinuses for sinusitis.

Fig. 4.5 Right orbital and characteristic saddle nose deformity in Wegener’s granulomatosis.
Treatment of AAV
(See BSPAR guidelines for treatments used in paediatric rheumatology, p 415.)

When considering therapy, it is useful to remember that most evidence for treatment is derived from adult trials. It is also useful to consider the different phases of the therapeutic journey for AAV:

- **The pre-diagnostic phase:** occasionally lasting years. Significant organ damage can accrue in this phase, or even death.
- **Induction of remission phase:** typically 3–6 months.
- **Maintenance of remission phase:** usually 18–24 months.
- **Therapy withdrawal phase:** not all patients achieve this.

The following general points are worthy of note:

- The key to successful treatment is early diagnosis to limit organ damage.
- Treatment for paediatric AAV is broadly similar to the approach used in adults and involves corticosteroids, cyclophosphamide, and in some individuals plasma exchange (particularly for pulmonary capillaritis and/or rapidly progressive glomerulonephritis—‘pulmonary-renal syndrome’) to induce remission; followed by low-dose corticosteroids and azathioprine to maintain remission.
- Antiplatelet doses of aspirin can also be considered empirically on the basis of the ↑ risk of thrombosis associated with the disease process.
- MTX in combination with corticosteroids may have a role for inducing remission in patients with limited WG.
- Co-trimoxazole is commonly added to therapeutic programmes for the treatment of WG, particularly in those with upper respiratory tract involvement, serving both as prophylaxis against opportunistic infection and as a possible disease-modifying agent.
- Newer immunosuppressive agents and immunomodulatory strategies such as MMF and rituximab (see the ‘RAVE’ and ‘RITUXIVAS’ trials in adults), have been reported to be effective at inducing or maintaining remission in adults with AAV and are increasingly used in children for recalcitrant disease.
- Anti-TNF therapy is less effective for the treatment of AAV, although has been used anecdotally in this context with some success in select patients.

Outcome of AAV

- The AAV still carry considerable disease-related morbidity and mortality, particularly due to progressive renal failure or aggressive respiratory involvement, and therapy-related complications, such as sepsis.
- The mortality for WG from one recent paediatric series was 12% over a 17yr period of study inclusion. The largest paediatric series of patients with WG reported 40% of cases with chronic renal impairment at 33 months of follow-up despite therapy.
- Mortality in paediatric patients with MPA during follow-up has been reported to be 0–14%.
- For CSS in children, the most recent series quotes a related mortality of 18%.

Further reading
Polyarteritis nodosa (PAN)

Background
- PAN is a necrotizing vasculitis associated with aneurysmal nodules along the walls of medium-sized muscular arteries.
- Despite some overlap with smaller-vessel disease, PAN appears to be a distinct entity and, in adults in Europe and the USA, has an estimated annual incidence of 2.0–9.0/million.
- Although comparatively rare in childhood, it is the most common form of systemic vasculitis after HSP and KD.
- Peak age of onset in childhood is 7–11yr, often with a ♀ preponderance.
- Classification criteria for PAN are not diagnostic criteria, and meeting classification criteria is not equivalent to making a diagnosis in an individual patient—see rest of section and Vasculitis classification, p 168.

Aetiology
- Unknown: possible interaction between infection and aberrant host response.
- There may be genetic factors that make individuals vulnerable to PAN and other vasculitides, but these are not yet defined.
  - There are reports of PAN occurring in siblings that add weight to this hypothesis, but there are no detailed genetic studies.
- There is a well-recognized association of PAN and familial Mediterranean fever in parts of the world where this is common.
- There are data to support roles for hepatitis B and reports of a higher frequency of exposure to parvovirus B19 and cytomegalovirus in PAN patients compared with control populations.
- HIV has also been implicated, and PAN-like illnesses have been reported in association with cancers and haematological malignancies. However, in childhood, associations between PAN and these infections or other conditions are rare.
- Bacterial superantigens may play a role in some cases.
- Occasional reports suggest immunization as a cause, but this is not proven.

Clinical features of PAN
A diagnosis is made by considering all clinical features in a patient, only some of which may be classification criteria. Clinical manifestations (and investigation findings) can be very confusing, especially in the early phase of the disease with absence of conclusive diagnostic evidence.
- The main systemic clinical features of PAN are malaise, fever, weight loss, skin rash, myalgia, abdominal pain, and arthropathy.
- Skin lesions are variable, and may masquerade as those of HSP or erythema multiforma. The cutaneous features described in a recent international classification exercise for PAN in children occurred commonly and were defined as follows:
  - Livedo reticularis—purplish reticular pattern usually irregularly distributed around subcutaneous fat lobules, often more prominent with cooling.
  - Skin nodules—tender subcutaneous nodules.
• **Superficial skin infarctions**—superficial skin ulcers (involving skin and superficial subcutaneous tissue) or other minor ischaemic changes (nailbed infarctions, splinter haemorrhages, digital pulp necrosis).

• **Deep skin infarctions**—deep skin ulcers (involving deep subcutaneous tissue and underlying structures), digital phalanx or other peripheral tissue (nose and ear tips) necrosis/gangrene.

• **Renal manifestations** such as haematuria, proteinuria, and hypertension.

• **GI features** and abdominal pain are relatively common and include:
  • Indeterminate intestinal inflammation: intestinal inflammation without characteristic histological features of either ulcerative colitis or Crohn’s disease. NB: routine mucosal gut biopsies rarely detect overt vasculitis since the small- and medium-sized arteries lie below the mucosa.
  • GI haemorrhage (upper and lower).
  • Intestinal perforation.
  • Panreatitis.

• **Neurological features** such as focal defects, hemiplegia, visual loss, mononeuritis multiplex; and organic psychosis may be present.

• **Other important clinical features** include: ischaemic heart and testicular pain. Rupture of arterial aneurysms can cause retroperitoneal and peritoneal bleeding, with perirenal haematoma being a recognized manifestation of this phenomenon, although this is rare.

**Differential diagnosis**

• Other 1st vasculitides: HSP, WG, MPA, KD. See relevant chapters, HSP p 179, WG p 188, MPA p 188, KD p 183.

• Autoimmune or autoinflammatory diseases:
  • JIA—particularly the systemic form.
  • JDM.
  • SLE.
  • Undifferentiated connective tissue disease.
  • Sarcoidosis.
  • Behçet’s disease.

• Infections:
  • Bacterial, particularly streptococcal infections, and sub-acute bacterial endocarditis.
  • Viral—many: specifically look for hepatitis B/C, CMV, EBV, parvovirus B19 and consider HIV.

• Malignancy: lymphoma, leukaemia, and other malignancies can mimic PAN.

**Diagnostic laboratory and radiological investigation**

**Blood tests**

• Anaemia, polymorphonuclear leucocytosis, thrombocytosis, ↑ ESR and CRP.

• Platelets are hyperaggregable.

• Circulating immune complexes or cryoglobulins may be present.

• Positive hepatitis B serology in children is unusual in association with PAN but can occur.
• ANCA are not thought to play a major part in the causality of PAN, but there are reports demonstrating their presence in some adults and children with PAN.
  • The presence of cytoplasmic ANCA (C-ANCA) with antibodies to proteinase 3 in a patient suspected of having PAN makes it mandatory to eliminate WG as a diagnosis.
  • Likewise, a significant titre of perinuclear ANCA (P-ANCA) with antibodies to myeloperoxidase would necessitate steps to eliminate microscopic polyangiitis (MPA) as the diagnosis.

**Tissue biopsy**
• Biopsy material is diagnostically important, especially skin or muscle, although tissue biopsy has overall low diagnostic sensitivity since the disease is patchy and vasculitis can be easily missed.
• The characteristic histopathological changes of PAN are fibrinoid necrosis of the walls of medium or small arteries, with a marked inflammatory response within or surrounding the vessel (Fig. 4.6).
• However, absence of such changes would not exclude the diagnosis, as the vasculitic features are variable and affected tissue may not have been sampled.
• Renal biopsy is usually not helpful and carries a greater risk than usual of bleeding and the formation of arteriovenous fistulae.

**Radiological tests**
• The most valuable investigative procedure is catheter-selective visceral digital subtraction arteriography to include flush aortogram and selective renal, hepatic, and mesenteric arteriography. This should be performed and interpreted only by those with expertise in this test in paediatric patients.
  • Arteriography findings include aneurysms, segmental narrowing, and variations in the calibre of arteries, together with pruning of the peripheral renal vascular tree (Fig. 4.7).
  • Treatment with prior corticosteroids will alter the arteriography and can result in false negatives.
  • Non-invasive arteriography such as CT or MR angiography (CTA/MRA) are not as sensitive as catheter arteriography for the detection of medium-sized vessel vasculitis such as PAN (discussed later in this list).
  • Consider formal cerebral arteriography if clinical and MRI features suggest cerebral vasculitis (see Cerebral vasculitis, p 209).
• Indirect evidence of the presence of medium-size artery vasculitis affecting renal arteries may be obtained by demonstrating patchy areas within the renal parenchyma of isotope uptake on Tc-99m dimercaptosuccinic acid (DMSA) scanning of the kidneys.
• Magnetic resonance angiography (MRA) usually fails to detect aneurysms of small- and medium-sized muscular arteries, although it may demonstrate large intra- and extrarenal aneurysms and stenoses/occlusions of the main renal arteries, and areas of ischaemia and infarction.
  • A caveat is that MRA may overestimate vascular stenotic lesions—CTA may also reveal larger aneurysms and arterial occlusive lesions.
and demonstrate areas of renal cortical ischaemia and infarction, but at the expense of high ionizing radiation exposure with less sensitivity than catheter arteriography.

- Echocardiography can be useful for the identification of pericarditis, valve insufficiency, myocarditis, or coronary artery abnormalities.

Fig. 4.6 (See also Colour plate 1.) PAN—skin biopsy.

Fig. 4.7 PAN—renal arteriogram.
Treatment
(See Chapter 3, The standard treatment of childhood vasculitis, p 174, for specific drug doses and protocols, and Chapter 2, BSPAR guidelines for treatments used in paediatric rheumatology, p 415.)

- In most patients, it is appropriate to treat aggressively to induce remission (typically 3–6 months), followed by less aggressive therapy to maintain remission (typically 18–24 months).
- In those presenting with mild predominantly cutaneous disease (see Chapter 2, Cutaneous PAN, p 198), corticosteroid alone may be appropriate, with careful monitoring of clinical and laboratory parameters as this is weaned.
- **Induction therapy:** high-dose corticosteroid with an additional cytotoxic agent such as cyclophosphamide:
  - Cyclophosphamide is usually given as pulsed monthly IV injections for up to 6 months or for shorter periods in children if remission is achieved.
  - Oral cyclophosphamide 2mg/kg per day for 2–3 months is an alternative, although for other vasculitides the IV regimen has been shown to have a more favourable therapeutic index.
- Aspirin 1–5mg/kg/day as an antiplatelet agent may be considered.
- **Maintenance therapy:** once remission is achieved, therapy with daily low-dose prednisolone and oral azathioprine is frequently used for up to 18–24 months.
  - Other maintenance agents include MTX, MMF, and ciclosporin.
  - Some advocate alternate day low-dose prednisolone in the maintenance phase with the intention of limiting steroid toxicity such as growth impairment although data to support this approach are limited.
- Adjunctive plasma exchange can be used in life-threatening situations (see Chapter 2, Vasculitis treatment, p 174).
- Biologic agents such as infliximab or rituximab have been used for those unresponsive to conventional therapy.
- Treatment response can be assessed using a modified Birmingham Vasculitis Activity Score (BVAS); the paediatric version of BVAS, or ‘PVAS’, is currently still being validated; and by monitoring of conventional acute phase reactants, urinary sediment, BP, and growth.

Outcome
- PAN, unlike some other vasculitides such as WG, appears to be a condition in which permanent remission can be achieved. Relapses can occur, but despite these, a real possibility of cure can be anticipated.
- However, if treatment is delayed or inadequate, life-threatening complications can occur due to the vasculitic process.
- Severe complications, especially infections, can occur from immunosuppressive treatment.
- In comparison with the almost 100% mortality rate in the pre-steroid era, mortality rates as low as 1.1% were reported in a recent retrospective multicentre analysis. However, this may not truly reflect mortality in circumstances of severe disease because 30% of patients in that series were considered to have predominantly cutaneous PAN.
- A mortality rate of 10% was recently recorded from a major tertiary referral centre seeing predominantly children with aggressive advanced disease.
• Late morbidity can occur years after childhood PAN from chronic vascular injury, possibly resulting in premature atherosclerosis. This remains a cause for concern and an area of ongoing research.

Further reading
CHAPTER 4 Systemic diseases

Cutaneous polyarteritis nodosa (cPAN)

Background and clinical features
- cPAN is a form of vasculitis affecting small- and medium-sized vessels limited to the skin.
- It is characterized by the presence of fever; subcutaneous nodular, painful, non-purpuric lesions with or without livedo reticularis occurring predominantly in the lower extremities; with no systemic involvement (except for myalgia, arthralgia, and non-erosive arthritis).
- In a recent international survey of childhood vasculitis, approximately 1/3 of children identified as having PAN were categorized as cPAN.
- The clinical course is characterized by periodic exacerbations and remissions that may persist for many years.
- Skin biopsy shows features identical to systemic PAN.
- ANCA are usually negative and the condition is often associated with serological or microbiological evidence of streptococcal infection.
- There is debate as to whether the condition should be classed as a separate entity or as a part of the spectrum of PAN since a proportion of cases appear to evolve into full-blown PAN.

Treatment of cPAN
- NSAIDs may suffice.
- Some require moderate doses of oral steroids.
- When streptococcal infection is implicated, penicillin may be effective.
  - Some recommend continuing prophylactic penicillin throughout childhood, as relapses are common and occur in up to 25% of cases in association with further streptococcal infections.
- When there is a lack of response to the above, or concerns about possible steroid toxicity, other agents may be considered:
  - IVIg has been successfully used.
  - Alternatives with anecdotal success for cPAN therapy include colchicine, hydroxychloroquine, azathioprine, MTX, dapsone (beware haemolytic anaemia as a relatively common and severe side effect of this agent), cyclophosphamide, and pentoxifylline.

Outcome of cPAN
- A minority of patients experience a persistence of cutaneous lesions through childhood.
- Overall it is uncommon for the condition to progress to PAN.
- However, it is mandatory for such patients to remain under surveillance to detect any evidence of developing systemic disease that would be an indication for intensification of treatment as per that of PAN.

Further reading
Takayasu arteritis

Background
- Takayasu arteritis (TA) is an idiopathic, chronic inflammation of the large vessels, affecting the aorta and its major branches.
- The disease is named after Mikito Takayasu, a Japanese ophthalmologist, who first described an association between retinal peri-papillary arterio-venous anastomoses and absent radial pulses.
- Other names include ‘pulseless disease’, aortic arch syndrome, or idiopathic aortoarteritis.
- Classification criteria are provided in the Vasculitis classification, p 168.

Epidemiology
- TA is more prevalent in Asian and African populations, and is rarer in Europe and North America. Most studies report an incidence of 1–3 per million/year in Caucasian populations—in Japan the estimated incidence is up to 100 times higher: 1 per 3000/year.
- In adult studies there is a 5:1 Q predominance. In children however gender ratios vary amongst different studies. A recent study from Southeast Asia and Africa report a Q:Q ratio of 2:1.
- TA is a rare vasculitis in children. Age of onset may range from infancy to middle age. The peak period of onset is in the 3rd decade of life.

Aetiopathogenesis
- The cause remains unknown.
- Genetic factors may play a role, and there are several reports of familial TA including in identical twins.
  - HLA associations include: HLA-A10, HLA-B5, HLA-Bw52, HLA-DR2, and HLADR4 in Japan and Korea; HLA B22 association has been described in the US population.
  - The presence of HLA Bw52 has been associated with coronary artery and myocardial involvement and worse prognosis.
- TA is described in association with RA, ulcerative colitis, and other auto-immune diseases suggesting an autoimmune mechanism for the pathogenesis of the disease.
- Circulating anti-aortic endothelial cell antibodies in patients with TA have been reported; their exact role however is yet to be determined.

Histopathology
- TA is characterized by granulomatous inflammation of all layers of the arterial vessels (panarteritis).
  - Inflammation of the tunica intima is followed by intimal hyperplasia leading to stenoses or occlusions.
  - Destruction of tunica elastica and muscularis cause dilatation and aneurysms.
  - Endothelial cell damage leads to a prothrombotic tendency.
  - The lesions have a patchy distribution.
  - The initial finding is neutrophil infiltration of the adventitia and cuffing of the vasa vasorum with proliferation and penetration of the latter within the tunica intima.
Various mixed chronic inflammatory cells including T cells contribute to granuloma formation in the tunica media and adventitia mediated by the release of interferon-γ and TNFα.

Later, the adventitia and media are replaced by fibrous sclerotic tissue and the intima undergoes acellular thickening, thus narrowing the vessel’s lumen and contributing to ischaemia.

In paediatric series:
- Occlusions and stenoses were present in 98% of the patients while aneurysms were only seen in 15.6% of the patients.
- Post-stenotic dilatations were present in 34% of cases.
- Lesions are most commonly seen in the subclavian arteries (90%), the common carotids (60%), the abdominal aorta (45%), the aortic arch (35%), and the renal arteries (35%); pulmonary arteries are involved in 25% of the cases.

Clinical features

Acute phase
Non-specific features of systemic inflammation (systemic, pre-stenotic phase). In children, up to 65% of TA present abruptly with systemic features:
- Pyrexia, malaise, weight loss, headache, arthralgias and/or myalgias.
- Rash (erythema nodosum, pyoderma gangrenosum).
- Arthritis.
- Myocarditis causing congestive heart failure (± hypertension) or valvular involvement (aortic valve most commonly affected followed by mitral valve).
- Myocardial infarction.
- Hypertension.
- Hypercoagulable state: thrombotic tendency.

Chronic phase
Features and signs 2° to vessel occlusion and ischaemia (stenotic phase):
- Asymmetric or absent pulses; a measured difference of >10mmHg on 4-limb BP monitoring is likely to indicate arterial occlusion.
- Systemic hypertension: commonest finding.
- Arterial bruits.
- Congestive heart failure 2° to hypertension and/or aortic regurgitation when the valve is affected.
- Angiodynia: localized tenderness on palpation of the affected arteries.
- Claudication.
- Coronary angina.
- Mesenteric angina presenting with abdominal pain and diarrhoea from malabsorption.
- Recurrent chest pain from chronic dissection of the thoracic aorta or pulmonary arteritis.
- Pulmonary hypertension.

CNS involvement
May be attributed to ischaemia ± hypertension: dizziness, or headache; seizures; transient ischaemic attacks, stroke.
Eye involvement
- Diplopia, blurry vision, amaurosis, visual field defect. Fundoscopy findings include:
  - Retinal haemorrhage
  - Micro aneurysms of the peripheral retina
  - Optic atrophy.

Renal involvement
- Renal hypertension due to renal artery stenosis with glomerular damage.
- Chronic renal failure.
- Amyloidosis.
- Glomerulonephritis (GN) has been described in association with TA: IgA nephropathy; membranoproliferative GN; crescentic GN; mesangioproliferative GN.

Differential diagnoses
- Other vasculitides including medium- and small-vessel vasculitis: Kawasaki disease; polyarteritis nodosa; Wegener’s granulomatosis is also recognized cause of aortitis.
- Infections:
  - Bacterial endocarditis
  - Septicaemia without true endocarditis
  - TB
  - Syphilis
  - HIV
  - Borelliosis (Lyme disease)
  - Brucellosis (very rare).
- Other autoimmune or autoinflammatory diseases: SLE; rheumatic fever; sarcoidosis; Blau’s syndrome.
- Non-inflammatory large vessel vasculopathy of congenital cause. Treatment with immunosuppression will be ineffective and could be harmful:
  - Fibromuscular dysplasia
  - William’s syndrome
  - Congenital coarctation of the aorta
  - Congenital mid-aortic syndrome
  - Ehler–Danlos type IV
  - Marfan syndrome
  - Neurofibromatosis type I.
- Other: post radiation therapy.

Laboratory investigations (also see Vasculitis investigation, p 172)
- Normochromic normocytic anaemia, leucocytosis, thrombocytosis; raised ESR, raised CRP—may not be present in chronic (stenotic) phase of illness.
- Elevated transaminases and hypoalbuminaemia.
- Deranged renal function tests in cases of renal involvement.
- Poikiloclonal hyperglobulinaemia.
Further tests required to exclude other causes mimicking TA or for disease monitoring

- Regular 4-limb BP measurement (preferably with a manual sphygmomanometer).
- In cases of significant peripheral artery stenosis, central BP measurements may be required.
- Renal function tests, urinalysis.
- Auto-immune screen.
- Baseline immunology tests including lymphocyte subsets, nitroblue tetrazolium (NBT) test.
- Blood cultures (acute phase).
- Mantoux test or interferon gamma releasing assays (IGRA).
- Syphilis serology.
- Tissue biopsy, rarely performed but should include microbiological culture, 16S and 18S ribosomal PCR if available to exclude bacterial and fungal infection respectively.

Imaging

- An echocardiogram (and ECG) and a CXR are simple 1st-line imaging tests and should be performed in all cases where TA is suspected.
- Conventional digital subtraction catheter arteriography is the method used routinely for obtaining a generalized arterial survey when TA is suspected, but essentially only provides ‘lumenography’ with no imaging of arterial wall pathology.
- MRI and MRA, and CTA, or a combination of these may help accurately diagnose TA and monitor disease activity, and (for MRA and CTA) provide cross-sectional aortic wall images allowing detection of arterial wall thickness and intramural inflammation.
  - MRI and MRA are gradually replacing conventional angiography in most centres and are useful for diagnosis and follow-up.
  - However, MR lacks sensitivity in evaluation of the distal aortic branches, and may overestimate the degree of arterial stenosis, especially in small children.
  - Cardiac MRI is increasingly employed to look for valvular involvement and/or myocarditis.
- Angiographic findings form the basis of one classification for TA (Takayasu Conference, 1994):
  - Type I. Classic pulseless type that affects blood vessels of aortic arch; involving the brachiocephalic trunk, carotid, and subclavian arteries.
  - Type II. Affects middle aorta (thoracic and abdominal aorta).
  - Type III. Affects aortic arch and abdominal aorta.
  - Type IV. Affects pulmonary artery in addition to any of the above types.
  - Type V. Includes patients with involvement of the coronary arteries.
- Doppler USS:
  - High resolution duplex US technology is a valuable tool in evaluation and follow-up of TA.
  - This modality offers high-resolution imaging of the vascular wall and can be useful for the detection of wall thickness.
• **18F-FDG-PET co-registered with CTA** can be a powerful technique combining information relating to the metabolic activity of the arterial wall (**18F-FDG** uptake detected using PET) with detailed lumenography (CTA) thus providing information on disease activity and anatomy. This technique is not available in all centres, and carries a high radiation exposure limiting its use for routine follow-up of disease activity.

**Diagnosis**

The diagnosis of TA is based on clinical and laboratory findings of systemic inflammation and/or of large-vessel ischaemia, raised and angiographic demonstration of lesions in the aorta or its major branches, with exclusion of other causes listed in the differential diagnosis.

**Treatment (also see [Vasculitis treatment, p 174](#))**

- Early diagnosis and aggressive treatment is fundamental for the outcome of the disease, although new lesions can continue to develop even in the presence of clinical remission in 60% of cases.
- Vascular damage already established in some patients will usually not respond to medical treatment.
- Medical management of TA includes: high-dose corticosteroids, usually in combination with MTX or cyclophosphamide for induction of remission. Maintenance agents include MTX, azathioprine, or more recently MMF. There are case reports of benefit with the use of infliximab.

**Hypertension**

- At least 40% of TA patients are hypertensive.
- Optimal control of hypertension is essential in the longer term since it is a major contributor to long-term morbidity.
- Medical treatment of hypertension in TA may be challenging since renovascular hypertension may not respond to medical therapy alone. **Seek specialist advice from a paediatric nephrologist.**
- Revascularization procedures may be required.

**Revascularization and other surgical procedures**

Techniques include:

- Angioplasty (including percutaneous transluminal angioplasty; or patch angioplasty), arterial bypass procedures, endarterectomy, arterial stenting, cardiac valve repair/replacement
- Surgery during the acute phase of the disease carries significant risk of re-occlusion and procedural complication, so should be deferred until the acute phase is treated
- These techniques should only be undertaken in centres with expertise.

Indications for revascularization include:

- Hypertension from stenotic coarctation of the aorta or renovascular disease.
- End-organ ischaemia or peripheral limb ischaemia.
- Cerebral ischaemia.
- Aortic or other arterial aneurysms, or aortic regurgitation.
CHAPTER 4 Systemic diseases

Prognosis

- There is usually a significant time lag (approximately 18 months, occasionally much longer) between initial presentation and diagnosis of TA in children. Arterial damage accrues during this pre-diagnostic phase, and influences prognosis.
- The course of the disease is variable, but most patients experience new lesions over time. Typically vascular inflammation persists even in patients thought to be clinically in remission.
- Aortic valve insufficiency and congestive heart failure are reported in 25%.
- Vascular claudication limiting activities occurs in up to 40%.
- Long-term mortality ranges from 10–30%:
  - The main causes of death include congestive cardiac failure, myocardial infarction, aneurysm rupture, or renal failure.
- After commencement of treatment approximately 60% will respond to corticosteroids while 40% will relapse when these are tapered off.
- Poor prognostic factors are severe aortic regurgitation, severe hypertension, cardiac failure, and aneurysms.

Further reading

Behçet’s disease

Behçet’s disease (BD) is a multisystem disease with a classic triad of recurrent oral aphthous ulcers, genital ulcers, and uveitis, which may also affect the skin, joints, GI tract, and CNS. Both small- and large-vessel vasculitis occurs, and in some patients there is a propensity to arterial and venous thromboses. Many now regard BD within the spectrum of autoinflammatory diseases.

Pathophysiology

- Unknown.
- BD has geographical variability, being more commonly found in the Mediterranean, Middle East, and Japan than in the USA and the UK.
- Likely autoinflammatory condition with possible environmental triggers in genetically susceptible individuals.
- HLA-B51 is significantly associated with BD.
- Recent studies also implicate the familial Mediterranean fever gene (MEFV) as an additional genetic susceptibility factor in some patients.
- Other genetic associations including MICA and TNF genes are thought to be the result of linkage disequilibrium with the HLA-B 51 gene, and are thus not causally associated.
- Pathergy: the skin hyper-reactivity response or pathergy test is not pathognomonic of BD but may be an important diagnostic indicator.
  Early or pathergy-induced cutaneous lesions in BD show a neutrophilic vascular reaction. This may be vascular or perivascular with a diffuse dermal neutrophilic infiltrate. Longstanding lesions may show a lymphocytic vasculitis. A typical pathergy test protocol is described later in this section.

Diagnostic criteria

The presence of recurrent oral aphthae (>3 times in one year), plus 2 of the following in the absence of other systemic diseases:
- Recurrent genital aphthae
- Eye lesions (uveitis or retinal vasculitis)
- Skin lesions
- Positive pathergy test.

Clinical features

Almost any organ can be involved due to the vasculitis affecting both arteries and veins of all sizes:
- Oral ulcers—painful, occur singly or in crops, affect lips, gingivae, buccal mucosa and tongue, last 1–2 weeks.
- Genital ulcers—affect scrotum in ♂, vulva in ♀, may result in scarring.
- Cutaneous lesions may be erythema nodosum-like, papulopustular, vesicular, abscesses, erythema multiforme-like eruptions, folliculitis, thrombophlebitis. Less commonly are pyoderma gangrenosum-type lesions, palpable purpura, purulent bullae, bullous necrotizing vasculitis and Sweet syndrome-like lesions.
- Ocular—iridocyclitis affecting the anterior segment or chorioretinitis, optic papillitis, retinal thrombophlebitis, arteritis.
- Arthralgia and arthritis usually affecting knees and ankles.
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- GI—nausea, vomiting, diarrhoea, weight loss. May mimic inflammatory bowel disease. Ulcers may occur anywhere but usually affect ileum and caecum.
- Neurological—headaches, paralysis, hyperparaesthesia, dementia, behavioural disorders, psychiatric problems, cerebellar signs, peripheral nerve palsies. Underlying pathology includes meningoencephalitis, cerebral venous thrombosis and ‘benign intracranial hypertension’, parenchymal inflammatory brain disease without obvious vasculitis (‘neuro-Behçet’s’), and true arterial vasculitis.
- Cardiovascular—myocarditis, pericarditis.
- Vascular—arterial or venous occlusions, varices, aneurysms.
- Respiratory—pulmonary infiltrates may be associated with pulmonary haemorrhage.
- Nephro-urological—haematuria and proteinuria (which may cause nephrotic syndrome), urethritis, orchitis, and epididymitis.
- Systemic—malaise, anorexia, weight loss.

Investigations

- FBC, CRP, ESR, routine clinical chemistry, thrombophilia screen (to exclude other causes of thrombosis), urinalysis.
- IgG, A, and M to exclude common variable immunodeficiency or other cause of low immunoglobulins (can present with oral ulcers).
- ANA, ds-DNA antibodies, RF, C3 and C4, ANCA, anticyclic citrullinated peptide antibodies, and antiphospholipid antibody are negative or normal but should be performed to exclude other autoimmune disease.
- Consider IgD and genetic testing for hyper IgD syndrome in atypical cases of suspected BD.
- Skin pathergy test—A 20-gauge needle is inserted obliquely 5mm into the skin of the anterior forearm. The presence of erythema, papules, erythematous papules, or pustules at 24–48h indicate a positive test result.
- MRI of the brain including MR arteriography and venography should be considered early for those with headaches to exclude neuro-BD or cerebral venous thrombosis.

Treatment

Evidence in children is limited. EULAR recommendations for treatment in adults are based on limited evidence and were not designed with children in mind, but provide a framework for the management of children. See Table 4.5.

General principles of treatment of BD in the young.

- Tailored to the individual and reflecting organ involvement and severity.
- Least toxic therapies should be tried first.
- Oral ulcers should be treated with topical agents before considering systemic drugs.
- Anti-TNFα agents (etanercept, infliximab, and adalimumab) are increasingly used before thalidomide in children.
Thalidomide is still used on a named-patient basis for children and adolescents with BD, typically for severe and resistant oral and/or genital ulceration, although it may also benefit other systemic symptoms of BD. Peripheral neuropathy and teratogenicity limit its use.

- Suggested dosing regimen: 0.5–1mg/kg (maximum 100mg) orally once a week at nighttime (to avoid daytime drowsiness). ↑ dose by adding another daily dose every week until symptoms controlled or until dose of 1mg/kg daily (whichever achieved first).
- Exclude pregnancy, and check peripheral nerve conduction studies prior to starting thereafter repeating nerve conduction 3–6-monthly (or after every 10g of total accumulative dose). Stop immediately if symptoms of neuropathy (e.g. numbness, tingling) develop.
- Remember that teratogenic risk also applies to mothers/female carers who handle the drug- full precautions always must be advised.
- The L isomer lenalidomide may be less toxic but have greater (or comparable) efficacy, but experience in paediatric BD is limited.

Prognosis—variable

- Eye disease can result in long-term visual impairment.
- Course may be worse in children than adults.
- O tend to be more severely affected.
- Reported mortality of 3%, usually due to vascular complications.

Table 4.5 EULAR 2008 recommendations for the treatment of Behçet’s disease"
### Chapter 4: Systemic diseases

#### Table 4.5 (Contd.)

<table>
<thead>
<tr>
<th>Recommendation</th>
<th>Category of evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>4. Anticoagulation:</strong> there are no controlled data on, or evidence of benefit from uncontrolled experience with anticoagulants, antiplatelet, or anti-fibrinolytic agents and management of deep vein thrombosis or for the use of anticoagulation for the arterial lesions of BD</td>
<td>IV</td>
</tr>
<tr>
<td><strong>5. Gastrointestinal involvement:</strong> there is no evidence-based treatment that can be recommended for the management of GI involvement of BD. Agents such as sulfasalazine, corticosteroids, azathioprine, TNFα antagonists and thalidomide should be tried first before surgery, except in emergencies</td>
<td>III</td>
</tr>
<tr>
<td><strong>6. Joint involvement:</strong> in most patients with BD, arthritis can be managed with colchicine</td>
<td>Ib</td>
</tr>
<tr>
<td><strong>7. Neurological involvement:</strong> there are no controlled data to guide the management of CNS involvement in BD. For parenchymal involvement agents to be tried may include corticosteroids, interferon-α, azathioprine, cyclophosphamide, MTX, and TNFα antagonists. For dural sinus thrombosis corticosteroids are recommended</td>
<td>III</td>
</tr>
<tr>
<td><strong>8. Ciclosporin neurotoxicity:</strong> ciclosporin should not be used in BD patients with CNS involvement unless necessary for intraocular inflammation</td>
<td>III</td>
</tr>
<tr>
<td><strong>9. Mucocutaneous involvement:</strong> the decision to treat skin and mucosal involvement will depend on the perceived severity by the doctor and patient. Mucocutaneous involvement should be treated according to the dominant or codominant lesions present. Topical measures (i.e. topical corticosteroids) should be the 1st-line treatment for isolated oral and genital ulcers. Acne like lesions are usually of cosmetic concern only. Thus, topical measures as used in acne vulgaris are sufficient. Colchicine should be preferred when the dominant lesion is erythema nodosum. Leg ulcers in BD might have different causes. Treatment should be planned accordingly. Azathioprine, IFNα and TNFα antagonists may be considered in resistant cases.</td>
<td>Ib</td>
</tr>
</tbody>
</table>

CNS, central nervous system; IFN, interferon; TNF, tumour necrosis factor.

**Categories of evidence:** Ia: meta-analysis of randomized controlled trials; Ib: randomized controlled trial; Iia: controlled study without randomization; IIb: quasi-experimental study; III: non-experimental descriptive studies such as comparative, correlation and case-control studies; IV: expert committee reports or opinion or clinical experience of respected authorities or both.

Central nervous system vasculitis in children

Background
- Central nervous system (CNS) vasculitis in children is an increasingly recognized inflammatory brain disease that continues to pose great diagnostic and therapeutic challenges. CNS vasculitis may occur as a primary disease that is isolated to the CNS (primary angiitis of the CNS, PACNS) or as a manifestation of an underlying systemic condition.
- The most common systemic inflammatory diseases and infections that may cause secondary CNS vasculitis are summarized in Box 4.1.

Diagnostic criteria
Diagnostic criteria for PACNS in adults were proposed by Calabrese et al. in 1992 and they include the following:
- An acquired neurological deficit.
- Angiographic and/or histopathological features of angiitis within the CNS, and
- No evidence of systemic condition associated with these findings.

Although a paediatric case definition of PACNS in children (cPACNS) has not been proposed, most reported cases fit the Calabrese et al. criteria.

cPACNS is broadly subdivided into two forms of the disease:
- Large–medium-vessel vasculitis (further divided into progressive and non-progressive, according to angiographic evidence of disease progression 3 months after diagnosis).
- Small-vessel vasculitis.

Epidemiology
The true incidence of cPACNS is difficult to establish as the condition is rare and there is lack of consensus on diagnostic criteria.

Clinical features
The clinical presentation of cPACNS is heterogeneous, with some children presenting with a rapidly progressive neurological deficit, whereas others have a slowly evolving disease course over weeks or months. Most common presenting features (in isolation or in various combinations) include the following:
- Acute severe headache.
- Focal neurological deficit.
- Gross motor deficit, hemiparesis.
- Cranial nerve involvement and optic neuritis.
- Concentration and cognitive deficits, behaviour, and personality changes.
- New onset of seizures.
- Acute loss of consciousness and symptoms of increased intracranial pressure caused by either intracerebral or subarachnoid haemorrhage, or a CNS mass lesion.
- Movement abnormalities.
- Constitutional symptoms (fever, fatigue, weight loss) are uncommon but can present in a minority of patients with small vessel vasculitis.
Box 4.1 Secondary central nervous system vasculitis in children

Inflammatory disorders
- Systemic lupus erythematosus
- Behçet’s disease
- Sjögren’s syndrome
- Juvenile dermatomyositis
- Scleroderma
- Inflammatory bowel disease.

Systemic vasculitides
- Polyarteritis nodosa
- Kawasaki disease
- Henoch–Schönlein purpura
- ANCA-associated vasculitides: Wegener’s granulomatosis; microscopic polyangiitis; Churg–Strauss syndrome.

Infectious/post-infectious
- Bacterial:
  - Streptococcus pneumoniae
  - Salmonella spp.
  - Mycoplasma pneumonia
  - Mycobacterium tuberculosis
  - Treponema pallidum
- Viral:
  - Hepatitis C virus
  - Cytomegalovirus
  - Epstein–Barr virus
  - HIV
  - Parvovirus B19
  - Varicella zoster virus
  - Enterovirus
  - Spirochete
  - Borrelia burgdorferi.
- Fungal:
  - Candida albicans
  - Aspergillus.

Other
- Malignancy
- Graft-versus-host disease.
Differential diagnosis
CNS vasculitis can be mimicked in both its clinical presentation and radiological manifestations by a number of inflammatory and non-inflammatory disorders, summarized in Box 4.2.

Investigations
The aim of the diagnostic work-up in patients with suspected cPACNS is to exclude causes of 2° CNS vasculitis or other neuro-inflammatory conditions that can present in the same way. In general there are no consistent or reliable laboratory abnormalities in children with cPACNS, and normal inflammatory markers by no means exclude an active vasculitic process in the CNS. The following investigations should be considered:

Laboratory investigations
- FBC and blood film, haemoglobin electrophoresis/sickle cell screen if patient is black or of Mediterranean ethnicity.
- ESR and CRP.
- ANA/dsDNA, ENA.
- ANCA.
- C3 and C4 levels.
- IgG, IgA, IgM.
- Clotting screen.
- Full thrombophilia screen including:
  - Lupus anticoagulant and anticardiolipin antibodies
  - Protein C
  - Protein S
  - Antithrombin
  - APC resistance ratio
  - Factor V Leiden
  - Methylenetetrahydrofolate reductase (MTHFR) and prothrombin G20210A gene mutations
  - Von Willebrand antigen levels may be elevated
  - CSF studies to establish cell count, protein levels, and exclude infection. CSF opening pressure should also be measured; CSF oligoclonal bands (simultaneous serum testing for total IgG required)
  - Plasma amino-acids and plasma homocysteine
  - Plasma lactate and ammonia
  - Alpha galactosidase A (to exclude Fabry’s disease)
  - Serology for mycoplasma, Borrelia burgdorferi, VZV.
- Ophthalmology assessment to look for retinal vasculitis, infection, or other inflammatory disease.
- Echocardiogram and ECG.

Imaging studies:
- MRI brain/spine and MRA.
- MRI in cPACNS reveals areas of acute ischemia in a vascular distribution when large–medium vessels are affected. In cases of small-vessel disease, the lesions may be multifocal and not necessarily conform to a specific vascular distribution.
- The parenchymal lesions may involve both grey and white matter, and meningeal enhancement has also been described.
Diffusion weighted imaging (DWI) identifies areas of ischaemia in large-vessel disease.

MRA provides an assessment of the vasculature and may reveal beading, tortuosity, stenosis, and occlusion of the vessels.

Conventional catheter arteriography (CA) continues to be the radiological gold-standard for identifying cerebrovascular changes in patients with suspected CNS vasculitis and is more sensitive than MRA at detecting distal lesions that affect small calibre vessels, and for lesions in the posterior part of the brain.

Brain biopsy for confirmation of vasculitis should be considered in difficult cases, but is rarely performed in children due to the invasiveness of the procedure. It should however be strongly considered in cases of high clinical suspicion but with negative arteriography findings or in cases of poor response to therapy.

Biopsy findings characteristically reveal segmental, non-granulomatous, intramural infiltration of arteries, arterioles, capillaries, or venules.

Non-lesional biopsy may be considered when lesions identified on imaging are not easily accessible.

**Treatment**

There are no randomized control trials (RCTs) to guide therapy.

Current therapeutic recommendations are based on those for systemic vasculitis (see *Vasculitis therapy*, p 174) and include:

- 6 months' induction therapy with IV cyclophosphamide: 500–1000mg/m² (max 1.2g) every 3–4 weeks (usually 7 doses); corticosteroids and antiplatelet doses of aspirin, followed by

- 1–2yr maintenance therapy with azathioprine (1.5–3mg/kg/day), low dose daily or alternate day corticosteroid, and continuation of aspirin.

MMF has also been reported to be effective in some cases to maintain remission.

Full anticoagulation may need to be considered on an individual patient basis.

Treatment of large-vessel non-progressive disease remains controversial. There may be a role for steroids and aspirin without cytotoxic immunosuppression.

**Outcome**

A recent study of 62 children with cPACNS suggested a poorer prognosis for patients presenting with: 1) a neurocognitive dysfunction, 2) multifocal parenchymal lesions on MRI, or 3) evidence of distal stenoses on arteriography.

Further long-term follow-up studies are necessary to accurately define the prognosis of this condition in children.
Box 4.2 Mimics of CNS vasculitis in children

- Arterial dissection
- Thromboembolic disease (congenital heart disease, inherited thrombophilia)
- Antiphospholipid syndrome
- Sickle cell disease
- Moyamoya disease (cerebral arteriopathy characterized by progressive steno-occlusive changes at the terminal portions of the bilateral internal carotid arteries with arterial collateral vessels at the base of the brain)
- Fibromuscular dysplasia
- Fabry’s disease
- Sneddon’s syndrome (morphologically fixed livedo reticularis and cerebrovascular accidents)
- CADASIL (cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy)
- Susac’s syndrome (acute encephalopathy, branch retinal artery occlusions and sensorineural hearing loss)
- Amyloid angiopathy
- Neurofibromatosis type I
- Hyperhomocysteinaemia
- Drug-exposure (cocaine, amphetamine, methylphenidate)
- Metabolic diseases:
  - Mitochondrial diseases
  - Leucodystrophies
  - Mucopolysaccharidoses
- Multiple sclerosis
- Acute disseminated encephalomyelitis (ADEM)
- Devic’s disease/neuromyelitis optica (CNS demyelinating condition affecting predominantly the spinal cord and optic nerves characterized by the presence of aquaporin-4 water channel IgG antibodies)
- Vitamin B12 deficiency
- Rasmussen syndrome (neurological disorder characterized intractable focal seizures, progressive hemiplegia and increasing cognitive impairment)
- Sarcoidosis
- Coeliac disease
- 1° and 2° haemophagocytic lymphohistiocytosis (HLH)
- Progressive multifocal leukoencephalopathy (JC virus)
- Lymphoma
- Glioma
- Migraine/vasospasm
- Post radiation therapy for CNS tumour.

Further reading
Other vasculitides

Background
The EULAR/PReS endorsed consensus criteria for the classification of childhood vasculitides include a category of "other vasculitides" for vasculitides where an aetiological process was defined, or in which no other classification category was appropriate (see p 168). These include:

- Behçet’s disease
- Vasculitis 2nd to infection (including hepatitis B-associated PAN), malignancies, and drugs, including hypersensitivity vasculitis
- Vasculitis associated with connective tissue diseases
- Cogan’s syndrome
- cPACNS
- Unclassified vasculitides.

Isolated cutaneous leucocytoclastic vasculitis, and hypocomplementic urticarial vasculitis are also described in this chapter. Behçet’s disease and cPACNS are described in their respective chapters.

Vasculitis secondary to infection, malignancies, and drugs, including hypersensitivity vasculitis

Vasculitis secondary to infection

- Many viruses (HIV, parvovirus B19, cytomegalovirus, varicella-zoster virus, and human T-cell lymphotropic virus- HTLV1) can be responsible for systemic vasculitis, the most frequent being hepatitis B virus-related polyarteritis nodosa (HBV-PAN), even though its incidence has decreased over the past few decades.
- Mixed cryoglobulinemia has been shown to be associated with hepatitis C virus (HCV) infection in adults, but has not been reported in children.
- Some bacteria, fungi, or parasites can also cause vasculitis, mainly by direct invasion of blood vessels or septic embolization.
- Effective antimicrobial drugs are mandatory to treat bacterial, parasitic or fungal infections, while the combination of antiviral agents (vidarabine, interferon-α) and plasma exchange has been proven to be effective against HBV-PAN.

Vasculitis secondary to drugs, including hypersensitivity vasculitis

- Therapeutic agents from virtually every pharmacological class have been implicated in the development of drug-induced vasculitis.
- Typically presents with cutaneous vasculitis alone (palpable purpura, macules, plaques, bullae and ulcers), low grade fever, arthralgia, and microscopic haematuria.
- More rarely can present with life-threatening systemic involvement, which may result in a severe and sometimes fatal illness.
- Withdrawal of the offending agent alone is often sufficient to induce prompt resolution of clinical manifestations.
- Steroids may be used for systemic involvement and azathioprine or other immunosuppressants may be appropriate for refractory disease.
**Vasculitis secondary to malignancy**
- In some patients, vasculitis occurs during the course of or prior to malignancies, most often haematological rather than solid tumours. Evidence of autoantibodies, immune complexes, and complement consumption is typically absent.
- Vasculitis may also occasionally be a complication of chemotherapy, radiation therapy, and bone marrow transplantation.

**Vasculitis associated with connective tissue diseases**
- Vasculitis associated with connective tissue disorders in children most commonly arises in the context of pre-existing SLE, 1st Sjögren’s syndrome (SS), systemic sclerosis, relapsing polychondritis, 1st antiphospholipid syndrome, juvenile dermatomyositis, or mixed connective tissue (see specific chapters on each of these entities, SLE p 223, Sjögren’s syndrome p 277, systemic sclerosis p 260, antiphospholipid syndrome p 279, mixed connective tissue or overlap syndromes p 275).
- The vasculitis may involve vessels of any size, but small-vessel involvement is predominant.
- Patterns of involvement vary with the associated underlying disorder, and range from isolated cutaneous involvement to life-threatening internal organ involvement.
- When vasculitis occurs in the setting of a pre-existing connective tissue disorder, it often correlates with disease severity and portends a poorer prognosis. Prompt recognition and treatment of vasculitis can dramatically improve the outcome for these patients.

**Isolated cutaneous leucocytoclastic vasculitis**
- This refers to cutaneous leucocytoclastic vasculitis without systemic vasculitis or glomerulonephritis.
- Histologically leucocytoclastic vasculitis appears as a neutrophil infiltration in and around small vessels, with neutrophil fragmentation (often referred to as ‘nuclear dust’ or leucocytoclasis), fibrin deposition, and endothelial cell necrosis.
- Treatment may require corticosteroids, colchicine, hydroxychloroquine, azathioprine, MTX or rarely dapsone (beware severe haemolytic anaemia and/or methaemoglobinaemia with dapsone).

**Hypocomplementaemic urticarial vasculitis syndrome**
- Hypocomplementaemic urticarial vasculitis syndrome (HUVS) is an uncommon immune complex–mediated entity characterized by urticaria with persistent acquired hypocomplementaemia.
- The disease is extremely rare in the paediatric population.
- In patients with HUVS, systemic findings include leucocytoclastic vasculitis, angio-oedema, laryngeal oedema, pulmonary involvement (interstitial lung disease, haemoptysis, pleural effusions), arthritis, arthralgia, glomerulonephritis, and uveitis.
- Laboratory findings include low levels of C1q, C2, C3, and C4. The binding of C1q antibodies to immune complexes is thought to be important in the pathogenesis of renal disease in HUVS.
- Treatment is individualized and is based on disease severity and will typically include corticosteroids and other immunosuppressive agents such as azathioprine or cyclophosphamide.
Patients may have significant morbidity and mortality, most commonly caused by chronic obstructive pulmonary disease and acute laryngeal oedema.

**Cogan’s syndrome**

- Cogan syndrome is a rare syndrome of:
  - Interstitial keratitis.
  - Vestibulocochlear symptoms (hearing loss and balance problems).
  - Occasionally aortitis.
- The cause is unknown, although autoantibodies and small-vessel vasculitis have been implicated.
- Typical Cogan’s is characterized by:
  - Ocular involvement: primarily interstitial keratitis and occasionally conjunctivitis, uveitis, or subconjunctival haemorrhage.
  - Audiovestibular involvement giving a clinical picture similar to Ménière’s disease accompanied with progressive hearing loss of hearing, usually ending in deafness within 1–3 months.
  - Other symptoms include fever, loss of weight, cardiac involvement (aortic insufficiency reported to up to 15% of patients), myalgia, arthralgia, mucocutaneous manifestations, GI and neurological involvement.
- The differential diagnosis is:
  - Susac syndrome (retinocochleocerebral vasculopathy, presenting with acute or subacute encephalopathy, branch retinal artery occlusions and sensorineural hearing loss as a result of small infarcts in the brain, retina and cochlea).
  - Other vasculitides.
  - Relapsing polychondritis.
  - SLE or Sjögren’s may cause similar symptoms.
- During attacks patients may have raised inflammatory markers. Detection of antibodies against corneal or inner ear antigens has been studied in small case series.
- Treatment includes corticosteroid therapy, while other immunosuppressants such as MTX, cyclophosphamide have been used with variable efficacy. Cochlear implantation may ultimately be necessary for hearing loss, and physiotherapy is required for the vestibular symptoms.
- The course of the disease is variable, with some patients experiencing episodes of ocular and audiovestibular symptoms at variable intervals with complete remission in between. In the longer term ~90% of patients suffer severe hearing loss while long-term ocular sequelae are rare. Systemic involvement is associated with the worst prognosis and can develop years after the initial onset of symptoms justifying prolonged close monitoring.

**Further reading**


Vasculitis mimics: non-inflamatory vasculopathies

Background
There are numerous non-inflamatory vasculopathies that may mimic the clinical, laboratory, radiological, and/or pathological features of the 1° vasculitides. Some will have associated musculoskeletal manifestations and will be referred to the paediatric rheumatologist. Awareness of these mimics is essential to avoid the use of unnecessary and potentially harmful immunosuppression and to direct management to the correct underlying cause of disease. The commonest of these vasculitis mimics are discussed in this section. Mimics of CNS vasculitis are covered in p 213.

Fibromuscular dysplasia (FMD)
- FMD is a non-inflamatory vasculopathy leading to stenoses of small- and medium-sized arteries, sometimes with poststenotic dilatation resembling aneurysms. It is a major differential diagnosis for Takayasu disease.
- FMD has been detected in almost every vascular bed although the most commonly affected are renal arteries (60–75%) followed by the cervico-cranial arteries (25–30%), non-renal visceral arteries (5%), and arteries in the extremities (5%). Intracranial arterial poststenotic dilatations have been reported in 7% of adult patients, but rarely in children.
- Patients can remain asymptomatic or present with signs of vascular insufficiency such as hypertension, stroke, abdominal pain, or claudication.
- The pathological classification for FMD is based on the arterial layer involved. Medial fibroplasia accounts for 80–90% of all cases and is characterized by a ‘string of beads’ appearance on angiography due to alternating areas of stenosis and aneurysmal dilatation involving the mid-to-distal portions of the vessel.
- It can be difficult to differentiate diffuse intimal FMD from large-vessel vasculitis, since their angiographic appearance can be similar. Histological examination may be required in these cases.
- Imaging investigations to be considered include:
  - Renal ultrasonography and Doppler studies.
  - CTA and MRA have been increasingly used for non-invasive imaging of the vascular tree.
  - However, selective catheter arteriography continues to be the radiological gold standard for delineating the extent of vascular involvement.
  - Cerebral perfusion scans or other measures to exclude cerebral arterial insufficiency should be carried out before angioplasty or surgical correction of renal artery stenosis to relieve hypertension, since a drop in BP can precipitate stroke in this situation.
- The management of hypertension associated with renal artery FMD involves antihypertensive therapy and revascularization with percutaneous angioplasty or other revascularization procedures for patients with significant renal artery stenosis, severe hypertension with
inadequate response, or intolerance of antihypertensive medication. Management of dissecting carotid artery due to FMD includes antiplatelet therapy and percutaneous angioplasty or surgical repair for patients with signs of cerebral vascular insufficiency.

**Vasculopathies involving the TGFβ-signalling pathway**

*Marfan syndrome*
- Incidence of Marfan syndrome is reported to be 1 in 10,000.
- Marfan syndrome results from mutations in the fibrillin-1 (*FBN1*) gene on chromosome 15, which encodes the glycoprotein fibrillin. Recent studies have suggested that abnormalities in the transforming growth factor-beta (TGFβ)-signalling pathway may represent a final common pathway for the development of the Marfan phenotype.
- Affected patients are usually taller and thinner than their family members. Their limbs are disproportionately long compared with the trunk (dolichostenomelia). Arachnodactyly, pectus excavatum, or carinatum are common features (Fig. 4.8).
- Ocular findings include ectopia lentis, flat cornea, cataract, glaucoma, retinal detachment.
- Cardiovascular involvement is the most serious complication associated with Marfan syndrome and comprises aortic root dilatation, aortic dissection involving the ascending aorta, and mitral valve prolapse.
- Patients can also present with spontaneous pneumothorax, stretch marks (striae atrophicae in the lower back), recurrent or incisional hernia, and dural ectasia.

*Management: seek expert cardiological advice*

The paediatric rheumatologist should be aware that evidence now suggests that the vasculopathy of Marfan syndrome is associated with dysregulation of TGFβ, and that this can be blocked using angiotensin II receptor 1 blockade (AT1 antagonists) such as losartan. Animal and human data have demonstrated that this can significantly slow the progression of aortic root dilatation and prolong life. Thus early referral to a paediatric cardiologist is essential. Other management of the vasculopathy under expert paediatric cardiology supervision may include:
- General measures: moderate restriction of physical activity, endocarditis prophylaxis, echocardiography at annual intervals.
- Beta-blocker therapy (propanolol 2–4mg/kg/day in divided doses) should be considered at any age if the aorta is dilated, but prophylactic treatment may be more effective in those with an aortic diameter of <4cm.
- ACE inhibitors (enalapril 0.08mg/kg/day—up to 5mg) reduce central arterial pressure and conduit arterial stiffness and may be useful.
- Prophylactic aortic root surgery should be considered when the aortic diameter at the sinus of Valsalva is >5cm.
Loeys–Dietz syndrome

• Loeys–Dietz syndrome is considered an autosomal dominant disorder associated with mutations in either of the TGFβ receptors (TGFβ R1/TGFβ R2).

• 2 subtypes of Loeys–Dietz syndrome have been identified:
  • Loeys–Dietz syndrome type I patients have both craniofacial and vascular disorders. The most characteristic craniofacial findings are hypertelorism and broad or bifid uvula or cleft palate, 2 of the 3 components of the clinical triad that also includes arterial aneurysms and tortuosity.
  • Loeys–Dietz syndrome type II patients may have a bifid uvula but do not have a cleft palate, craniosynostosis, or hypertelorism.

• Additional manifestations include blue sclera, malar hypoplasia, exotropia, and retrognathia. Cervical spine instability, pectus deformity, arachnodactyly, craniosynostosis, scoliosis, and joint laxity are some of the many musculoskeletal manifestations. Patients may also have congenital cardiac anomalies (bicuspid aortic valve).

• Due to the high risk of death from aortic aneurysm rupture, patients should be followed closely to monitor aneurysm formation, which can then be corrected with vascular surgery.

• The role of TGFβ antagonism is currently being explored in this condition too.

Fig. 4.8 Pectus excavatum of moderate severity.
CHAPTER 4  Systemic diseases

Ehler–Danlos syndrome (vasculopathic EDS) type IV
- EDS type IV is an autosomal dominant disorder that results from mutations in the type III pro-collagen gene (*COL3A1*).
- Although joint and skin laxity is not common, patients often have easy bruising, thin skin with visible veins, or characteristic facial features (loss of subcutaneous fat and collagen, referred to as acrogeria).
- Arterial complications may include aortic or other arterial aneurysm, dissection, and carotico-cavernous sinus fistulae. Patients can present acutely with life-threatening rupture of the intestines, the gravid uterus, or other viscera.
- There is currently no preventative treatment for this condition but close follow-up to monitor aneurysm formation is recommended.

Grange syndrome
- Grange syndrome is an hereditary disorder that is associated with a variable combination of multiple arterial stenoses and aneurysms, brachydactyly, and syndactyly of the hands and feet, bone fragility consistent with a mild form of osteogenesis imperfecta, learning disability, and cardiac defects (PDA, VSD, bicuspid aortic valve).
- No underlying genetic defect has yet been determined.

Susac’s syndrome
- Susac’s syndrome is a microangiopathy of unclear aetiology that is more frequently reported in young adult females.
- Susac’s syndrome is characterized by the typical clinical triad of acute or subacute encephalopathy, branch retinal artery occlusions, and sensorineural hearing loss that results from small infarcts in the brain, retina, and cochlea.
- Typical findings on brain MRI are multiple small white-matter hyperintensities; grey-matter involvement may also be seen. CSF analysis usually reveals a lymphocytic pleocytosis and elevated protein levels. Histological evidence of microangiopathic infarct is seen, without evidence of vasculitis or thrombosis.

Degos disease (DD)
- DD, also known as malignant atrophic papulosis, is characterized by thrombo-occlusive vasculopathy affecting the skin and various internal organs.
- DD has been described in 2 forms: the limited benign cutaneous and the lethal multiorgan systemic variant.
- In the skin, DD initially manifests with erythematous, pink or red papules that leave scars with pathognomonic, central, porcelain white atrophic centres.
- In the systemic form, GI involvement has been reported in the majority of patients and may manifest as abdominal pain, GI bleeding or bowel perforation. Central and peripheral nervous system, heart, lung, eye, pancreas, adrenal gland, and kidney involvement have been described.
- There has been no proven effective treatment for DD. Antiplatelet agents including aspirin and dipyridamole have been reported to reduce the formation of new skin lesions but there is no evidence in their role to prevent systemic complications. Immunosuppressants, anticoagulants,
and plasma exchange have been shown to be ineffective. Since corticosteroids may worsen other forms of occlusive vasculopathy, they should be used with caution in DD. Prompt surgical intervention is often needed for bowel infarction, perforation, or intracranial haemorrhage.

- The systemic form has a poorer prognosis and is usually fatal within the first 2yr after diagnosis because of major organ involvement.

**Livedoid vasculopathy (LV)**

- LV is an occlusive vasculopathy characterized by thrombosis and ulceration of the lower extremities.
- While the etiology of LV remains unclear, it likely has a prothrombotic pathogenesis. Factor V Leiden mutation, heterozygous protein C deficiency, and hyperhomocysteinaemia have been associated with LV. In addition, plasminogen activator inhibitor (PAI)–1 promoter 4G/4G genotype has also been linked to the disease.
- Skin biopsies reveal segmental hyalinizing vascular involvement of thickened dermal blood vessels, endothelial proliferation, and focal thrombosis without nuclear dust. No true vasculitis is evident.
- The initial clinical findings are typically painful purpuric macules or papules on the ankles and the adjacent dorsum of the feet. Patients may have a history of livedo reticularis on their lower legs. The initial lesions, which often appear in clusters or groups, eventually ulcerate over a period of months and years and form irregular patterns of superficial ulcers. When the ulcers finally heal, they leave behind atrophic porcelain-white scars, which are referred to as ‘atrophie blanche’.
- Small- and medium-sized vasculitides, such as isolated cutaneous leucocytoclastic vasculitis and PAN occasionally present with ulceration resulting in ivory-white, stellate scarring on the lower limbs and may be difficult to differentiate from LV.
- A number of therapies have been employed with variable effect:
  - Corticosteroids in combination with pentoxifylline have been used in cases of widespread LV. Pentoxifylline is believed to enhance the blood flow in the capillaries, making red blood cells more flexible and thereby reducing viscosity effectively. **However, anecdotaly some report that corticosteroids can worsen the occlusive vasculopathy associated with LV.**
  - As thrombogenic mechanisms may be involved in the disease pathogenesis, anticoagulant therapy (low-molecular-weight heparin, warfarin) and aspirin are often tried, but with variable efficacy.
  - Hyperbaric oxygen therapy has been used in intractable cases of LV with good effect in some cases.
Pseudoxanthoma elasticum (PXE)
- PXE is a rare, genetic disorder characterized by progressive calcification and fragmentation of elastic fibres in the skin, the retina, and the cardiovascular system, which is termed as elastorrhexia.
- PXE is caused by mutations in the ATP-binding cassette transporter C6 (ABCC6) also known as multidrug resistance-associated protein 6 (MRP6) gene.
- Patients present with:
  - Characteristic skin lesions that can appear during childhood. These are small, yellow papules of 1–5mm in diameter in a linear or reticular pattern which may coalesce to form plaques. The skin has a cobblestone-like appearance. These skin changes are first noted on the lateral part of the neck and later can involve any part of the body.
  - Ocular manifestations: angioid streaks of the retina, which are slate grey to reddish brown curvilinear bands radiating from the optic disc.
  - Cardiovascular manifestations include: calcification of the elastica media and intima of the blood vessels leading to a variety of physical findings, mitral valve prolapse.
  - Renal artery involvement leads to hypertension.
  - Mucosal involvement leading to GI haemorrhage.
- There is currently no treatment available for PXE.

Further reading