Emergency situations in rheumatology with a focus on systemic autoimmune diseases

Jiri Vymetal, Martina Skacelova, Andrea Smrzova, Anna Klicova, Marketa Schubertova, Pavel Horak, Josef Zadrazil

Background and Aim. Rheumatic diseases are commonly considered chronic conditions. However, acute manifestations can be very severe and represent a diagnostic problem. Examples are systemic lupus erythematosus with acute flare, glomerulonephritis, CNS disorders and catastrophic antiphospholipid syndrome, scleroderma with interstitial lung disease, pulmonary hypertension and renal crisis and polyangiitis with alveolar haemorhage and acute respiratory failure. This aim of this paper is to overview emergency situations which can be encountered in the care of patients with autoimmune systemic diseases and vasculitides.

Methods. A Pubmed search for both original and review articles, recent textbooks and current guidelines related to rheumatic diseases with possible acute situations were included in this review article. Relevant image documentation was obtained at the site over the past several years of observation.

Conclusions. This paper provides an overview of facts and emergency situations which can be encountered in the care of patients with autoimmune systemic diseases and vasculitides. It is directed at clinicians working in intensive care. It provides a differential diagnostic overview and information which is rare and commonly underestimated.

Key words: SLE, scleroderma, polyangiitis, dermatomyositis, emergent

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INTRODUCTION

Rheumatic diseases are in general chronic, slowly developing and progressing illnesses. Nevertheless, acute flares and corresponding symptoms are not infrequent, and can reflect an emergency and life-threatening event, requiring urgent intervention and intensive care. Patients with rheumatic disease may be referred to different medical departments (e.g. rheumatology, internal medicine, neurology, or cardiology) according to predominant symptom(s). The complexity of systemic disease signs and symptoms requires, in the case of life-threatening events/complications, multidisciplinary cooperation, including rheumatologist’s consultancy, and immediate patient’s admission to the ICU, to ensure adequate comprehensive intensive care.

Covered areas

The list of rheumatic diseases and related health statuses could be an extensive one. In this article, however, we shall focus on diseases with major clinical impact, i.e. systemic connective tissue disorders and vessel inflammatory disorders (vasculitis). Apart from basic clinical definitions and a general description of the diseases, we focus on the disease specific, life-threatening manifestations. Table 1 provides an overview of the field.

SYSTEMIC LUPUS ERYTHEMATOSUS (SLE)

Definition

SLE is an autoimmune inflammatory disease, predominantly affecting fertile women. Hyperactivity of B-cells is characteristic of the disease, together with overproduction of organ nonspecific autoantibodies that may participate on immunocomplexes formation. Deposition of immunocomplexes leads to tissue inflammatory impairment. Given that SLE is a highly heterogenic disorder, its clinical picture is rather assorted. In general, SLE can be split into several, clinically and laboratory defined subtypes.

Signs and symptoms

Fever, fatigue and weight loss are frequent symptoms of an acute flare of SLE. The course of the disease is characterized by remissions and exacerbations. Production of organ nonspecific antibodies against nuclear, cytoplasmic as well as cell surface self-antigens is one of the typical characteristics of the disease. Most frequent clinical symptoms include skin, joints, cardiovascular and renal disorders, as well as impaired respiratory and nervous systems and hemopoesis. Active SLE may end-up in the afflicted organ failure. Severe forms of the disease are associated with marked mortality.
Life-threatening disorders in SLE

Heart and cardiovascular system impairment

Cardiac impairment is quite frequent in SLE. In general, there are four major forms:

Valve impairment (Valve disease)

Mitral regurgitation is the most frequent event, usually hemodynamically insignificant. Valvular vegetation of variable size, from small nodules to large verrucous vegetation (Libman-Sacks endocarditis), are the most frequent cause of the valve dysfunction and/or insufficiency (mitral valve insufficiency is more frequent than aortal valve insufficiency) (ref. 1).

Pericardial disorders

Pericarditis, together with pleuritis and diffuse peritonitis are classified within the group of serositis. Pericarditis, which is often asymptomatic, frequently associates with other serositis in active SLE. Sharp and stabbing pain localized behind the lower part of the breastbone (sternum) is a typical symptom, frequently coupled with pleural friction rub. Large volume pericarditis, with high risk of cardiac (pericardial) tamponade is rare.

Myocardial dysfunction

Signs of heart failure, tachycardia or arrhythmia may be reflections of a myocardial dysfunction, accompanied by cardiomegaly. Diastolic impairment is the most frequent finding on echocardiogram, however, dilatation and diffuse hypokinesia with depression of the left ventricle systolic function may be found too. Acute myocarditis often goes together with pericarditis.

Ischemic heart disease

Cardiovascular mortality related to SLE reflects one of the highest mortality figures, out of the total cardiovascular mortality. The myocardial infarction relative risk

Table 1. Acute disorders in systemic rheumatic diseases.

<table>
<thead>
<tr>
<th>Type of disease</th>
<th>Possible related acute disorder</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rheumatoid arthritis</td>
<td>atlantoaxial dislocation, scleromalacia perforans, vasculitis</td>
</tr>
<tr>
<td>SLE (Systemic lupus erythematosus)</td>
<td>seizures, psychosis, encephalopathy, pericarditis, myocarditis, endocarditis, pneumonitis, ARDS, acute glomerulonephritis, hypertensive crisis, acute pancreatitis, polyserositis, infection/sepsis</td>
</tr>
<tr>
<td>Antiphospholipid syndrome</td>
<td>acute cerebrovascular event, acute myocardial infarction, retinal vascular thrombosis, pulmonary embolism and infarction, placental ischemia, spontaneous abortion, catastrophic antiphospholipid syndrome</td>
</tr>
<tr>
<td>Scleroderma and mixed connective tissue disease (MCTD, Sharp’s syndrome)</td>
<td>renal crisis, right-sided heart failure, digital ischemia</td>
</tr>
<tr>
<td>Dermatomyositis</td>
<td>acute lung disease, respiratory failure, bowel perforation</td>
</tr>
<tr>
<td>Vasculitis</td>
<td>pulmonary-renal syndrome, rapidly progressive glomerulonephritis, acute renal failure, acute lung disease, respiratory failure, pancreatitis, encephalopathy, arterial and venous thrombotic complications, necrosis</td>
</tr>
<tr>
<td>Parainfectious manifestations</td>
<td>reactive arthritis, septic arthritis, septic spondylodiscitis, pyogenic myositis</td>
</tr>
</tbody>
</table>

MCTD - mixed connective tissue disorder, ARDS - acute respiratory distress syndrome
in SLE corresponds to 2.27; the relative risk for CVA (stroke) equals to 2.05 and the relative risk for atherosclerosis rises to 7.1. Atherosclerosis genesis is related to a higher presence of traditional risk factors (obesity, smoking, metabolic syndrome), however, other factors specific for active SLE are involved in its subsequent development (disease activity, antiphospholipid antibodies, immunocomplexes, drug related adverse effects) (ref.4).

Acute lung disease

Lupus pneumonitis is a rare manifestation of active SLE disease. Its symptoms include fever, breathlessness, productive cough (with small quantity of sputum), tachypnea, hemoptysis, pleuritic pain and hypoxemia. Physical examination will display bilateral basal crepitus, and central cyanosis in more severe cases. Bilateral patchy alveolar infiltrates may be found on chest X-ray. Therefore, an infection as a potential option must be diagnostically excluded.

Pulmonary hemorrhage is a rare6, but serious and frequently fatal event. It shows clinical course similar to lupus pneumonitis, with rapid progression and deterioration of patient health status, which causes significant patient anxiety. Tachypnea, hypoxemia and tachycardia are regularly found, and ARDS can easily evolve (Fig. 1). Intraalveolar or external hemorrhage can be present, with frequent and rapid hemoglobin level reduction. Histopathology provides evidence of intraalveolar bleeding, with intact erythrocytes and macrophages packed with hemosiderin. Vasculitis may not be exhibited; nonetheless, microangiitis has been described from time to time, which corresponds to inflammatory necrosis of alveolar capillaries and small muscular arterioles.

Acute reversible hypoxemia: Its pathogenesis remains unclear. Oxygenation capacity of the lung may be reduced due to neutrophil aggregation within the pulmonary vessels. Inside neutrophils we can see residues of the complement decay7,8.

Central Nervous System Impairment – Neuropsychiatric and neuropsychological Symptoms

The prevalence of neuropsychiatric symptoms in SLE, as reported in clinical trials, is rather heterogeneous, ranging from 10% in early stages up to 80% during the later course of the disease. This heterogeneity reflects different criteria for evaluation of CNS impairment in SLE. The American College of Rheumatology (ACR) has defined the standards for diagnosis of 19 neuropsychiatric symptoms accompanying SLE. Detailed description of the standards is available on: www.rheumatology.org/publications/ar/1999/aprilappendix.asp?aud=mem.

Cerebral infarction

The risk of cerebral infarction and sudden death is significantly increased for patients with SLE. The cerebral infarction may be presented either as a classical stroke or as a transitory ischemic attack. The recurrence risk is high. There is a strong association between these events and antiphospholipid antibodies9,10. Patients with antiphospholipid syndrome usually suffer from repeating transitory ischemic events. These can lead to a significant cognitive disorder.

Seizures

Generalized or partial seizures may be the primary symptom of the disease. Their etiopathogenesis is diverse, could be the inflammatory damage of the CNS or the presence of an "old” scar. The presence of the antiphospholipid antibodies may increase the risk of seizures, as well as hypertension, infection, tumors, injuries, ischemic cerebral events, vasculopathies or drug toxicity (e.g. high doses of antimalarial) can do11. Seizures with no clear local etiology (negative findings on angiogram, CT scan or MRI) probably result from existing vasculopathy.

Acute confusional episodes

In SLE these are multifactorial, partly due to the disease activity, partly due to metabolic disorders and/or therapy. The status reflects the complexity of the disorder, with reduced ability to maintain attention and/or concentration. Behavioral and mood disorders, psychomotor restlessness and different levels of consciousness disorders may accompany episodes. These delirious conditions, developing within few hours or days, may end up in coma.

Renal impairment

Renal impairment per se and/or in combination with medication-induced nephrotoxicity may significantly increase both morbidity and mortality in patients with SLE. Appropriate diagnosis is based on detailed clinical and laboratory examinations, with supplementary histological examination of renal biopsy, if needed.

Glomerulonephritis is the fundamental pathogenic factor of renal impairment in SLE. According to WHO classification (revision 4), there are 6 basic histological classes of glomerulonephritis, which currently have been amended by activity and chronicity indexes. Lupus nephritis is often diagnosed in younger patients with symptomatic SLE. The degree of renal insufficiency and nephritis related proteinuria can vary individually.

Vascular pathology Capillary microthrombi or thrombosis of renal vessels are the major cause of the acute renal failure in the context of SLE complication and antiphospholipid syndrome.

Acute interstitial nephritis is another potential manifestation of the disease. It is associated with immunocomplex deposits distributed along the tubular basal membrane. Both proximal and distal tubular syndromes are present in many SLE patients, and frequently linked to manifestations of secondary Sjögren’s syndrome and positivity of anti-Ro (SSA) and/or anti-La (SSB) antibodies. This type of nephritis does not correlate with the existence and activity of glomerulonephritis.

Pregnancy and Lupus Erythematosus

High disease activity before conception and preexisting renal impairment are major risk factors for the disease flare in the course of pregnancy or after its termination. The risk of flare has been partially reduced by comprehen-
sive medical care and systematic monitoring of the SLE patients. On the other hand, the first manifestation of lupus during pregnancy or immediately after delivery may develop as a serious, even catastrophic case. The disease should be considered always in case of any relevant clinical symptomatology appears in the course of pregnancy ("butterfly" rash or other skin symptoms, alopecia, arthritis, proteinuria with active sediment, psychosis, chorea, pleurocarditis, vasculitis, etc.). Women with SLE have increased risk of preeclampsy and hepatopathy.17, 18.

Diagnosis
The disease diagnosis is based on the appropriate interpretation of history, clinical, laboratory and paraclinical findings. Determination of positivity of antinuclear autoantibodies (ANA), anti-dsDNA antibodies, ENA antibodies and antiphospholipid antibodies or reduced levels of complement play a crucial role in the disease diagnosis.

It is important, within the context of the SLE related life-threatening events, to rule out any other systemic connective tissue disorder and/or vasculitis, and particularly any potential infectious complication in differential diagnosis. Infections related to SLE often have an insidious course, which can be further altered by prescribed immunosuppressive therapy.

Therapy
Treatment of SLE is focused on management of the disease activity, maintenance of remission, prevention of severe organ impairment, and prevention or management of secondary complications. The treatment strategy is highly individual. Patients in advanced stage of the disease, with multiorgan failure should be managed first with the aim of saving their lives. In the stable disease, it is important to evaluate the disease activity, extent of organ impairment, quality of life, and potential toxicity of intended or already prescribed medication. It is important to mention, that many drugs commonly used in SLE are "off label". This concerns not only many of immunosuppressive drugs but also for example rituximab.

Table 2 summarizes simplified recommendations for the therapy of SLE, according to a type and severity of the disease.19-21.

Table 2. SLE therapy options, according to stage and severity.19-21.

<table>
<thead>
<tr>
<th>Stage and severity</th>
<th>Induction therapy</th>
<th>Maintenance therapy</th>
<th>Second line therapy (previous therapy failure or intolerance)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dermo-articular form</td>
<td>Hydroxychloroquine 200-400 mg p.o. (≤ 10mg/day or 0.125 mg/kg EOD)</td>
<td>Hydroxychloroquine 200-400 mg p.o.</td>
<td>MTX (7.5-25 mg/week)</td>
</tr>
<tr>
<td>Mild organ impairment</td>
<td>High dose prednisone (0.5 - 1.0 mg/kg/day) - with stepping down ± AZA (1-3 mg/kg/day); for 3-6 months</td>
<td>Low dose prednisone (≤ 10 mg/day or 0.125 mg/kg EOD) ± AZA (1-2 mg/kg/day) Hydroxychloroquine 200-400 mg p.o.</td>
<td>MTX (7.5-25 mg /week)</td>
</tr>
<tr>
<td>Moderate organ impairment</td>
<td>High dose prednisone (0.5 - 1.0 mg/kg/day) - with stepping down + MMF (2 g/day) or AZA (2-3 mg/kg/day) for 6-12 months, or 6 pulses of CFA (once a month)</td>
<td>Glucocorticoids + AZA (1-2 mg/kg/day) or MMF (1-1.5 g/day) Hydroxychloroquine 200-400 mg p.o.</td>
<td>MTX (7.5-25 mg /week)</td>
</tr>
<tr>
<td>Severe organ impairment</td>
<td>Pulses of methylprednisolone + CFA (Once a month) + Prednisone 0.5 mg/kg/day p.o. with stepping down, for 6-12 months</td>
<td>Glucocorticoids + AZA (1-2 mg/kg/day) or MMF (1-2 g/day) Hydroxychloroquine 200-400 mg p.o.</td>
<td>MMF 2.3 g/day and/or Rituximab</td>
</tr>
</tbody>
</table>

AZA - azathioprine; CFA - cyclophosphamide; MMF - mycophenolate mofetil; MTX - methotrexate;
ANTIPHOSPHOLIPID SYNDROME

Definition
Antiphospholipid syndrome (APS) is represented by symptoms related to the presence of antiphospholipid antibodies (lupus anti-coagulant, anti-cardiolipin anti-bodies, anti-beta 2 glycoprotein anti-bodies). Antiphospholipid syndrome can exist per se, or it may accompany other systemic connective tissue disorders, especially SLE, as so-called secondary APS (ref.23).

Signs and symptoms
The coagulation disorders, associated with laboratory pathology (prolonged aPTT, with correction by plasma of a healthy donor; false positive Wassermann reaction), together with arterial and/or venous thrombosis and organ impairment are predominant symptoms of APS. Thrombocytopenia and other associated symptoms are frequently present. Stillbirth and spontaneous abortions may be pregnancy related APS symptoms.

Life-threatening disorders
Life-threatening disorders related to APS are mainly vascular thrombosis, pulmonary embolism, arterial occlusions, cerebrovascular event, and myocardial infarction or microangiopathic hemolysis. Preeclampsia, eclampsia and HELLP syndrome are potentially fatal APS related disorders connected with pregnancy. These are associated with a high risk of stillbirth or spontaneous abortion24. The catastrophic antiphospholipid syndrome is of special interest. It represents a rare manifestation of the disease, with a vast thrombotic disorder and multiorgan failure. It may be fatal, in association with the development of disseminated intravascular coagulation, with increased levels of fibrin degradation products, D-dimers and fibrinogen consumption25. Clinical criteria of catastrophic antiphospholipid syndrome include:
- History of APS or presence of antiphospholipid antibodies
- Three or more episodes of organ thrombosis, within the last week
- Biopsy confirmation of microthrombi
- Exclusion of other reasons for organ thrombosis and microthrombi

Therapy
Therapy of primary and secondary APS is almost the same. In secondary APS, immunosuppression may be necessary to reduce the activity of the underlying basic pathology. Heparin (LMWH or unfractionated), warfarin and acetylsalicylic acid (ASA) are the most frequently used medication. Hydroxychloroquine may be beneficial for patients with SLE-APS. Treatment of APS related microangiopathy with hemolysis is similar to the treatment for thrombotic thrombocytopenic purpura/hemolytic-uremic syndrome (TTP/HUS). Apart from symptomatic treatment, it includes repeated plasmapheresis in addition. The catastrophic APS therapy is based on elimination of potential stir-up infectious factors and combined medication with antiagulants (heparin, LMWH) and high dose methylprednisolone, together with repeated plasmapheresis, in case of microangiopathy. Daily intravenous immunoglobulins, in a dose of 400 mg per kilogram of body weight, may serve as add-on therapy. How many plasmapheresis are needed depends on clinical response. According to some reports, anti-cardiolipin anti-bodies reduction of 95% can be achieved after the fifth plasmapheresis. Use of rituximab may be beneficial too26.

Prognosis
If not recognized early enough, APS, especially its catastrophic form, may be fatal, with a high rate of mortality.

SYSTEMIC SCLERODERMA (SSc)

Definition
Systemic sclerosis or systemic scleroderma is a chronic connective tissue disease. CREST syndrome, also known as the limited cutaneous form of systemic sclerosis (Ic-SSc), is a multisystem connective tissue disorder, which is limited mainly to the skin on the face, hands and feet. The acronym "CREST" refers to the five main features: Calcinosis, Raynaud’s phenomenon, esophageal dysmotility, sclerodactyly and telangiectasia. Diffuse cutaneous scleroderma covers most of the skin, and is at risk of progressing to the visceral organs, including the kidneys, heart, lungs and gastrointestinal tract. The disease is characterized by fibrosing sclerosis of peripheral and visceral vessels, over production of collagen in connective tissues, changes in macro vascularity and by humoral and cellular immunity disorders.

Signs and Symptoms
Raynaud’s phenomenon (RF) together with trophic changes and hardening of the skin are predominant symptoms of the disease (Fig. 2). Impairment of gastrointestinal tract, lung, heart and kidney may be present too, depending on the stage of the disease27.

Life threatening manifestation of SSc

Renal crisis
Renal impairment is a frequent symptom in SSc, mostly as mild renal insufficiency with a good prognosis28. On the other hand, severe renal crisis is an acute, serious and life-threatening event, with an incidence of 8-10% among patients with limited SSc and 10-20% in patients with diffuse form of the disease29. Usually, it develops within the first five years of onset of the disease. Renal crisis reflects thrombotic microangiopathy of kidney, and is similar to TTP/HUS or APS. The histology displays proliferation and thickening of arcuate and interlobular arteriole intima, resulting in narrowing or full obliteration of vessels. There appears the typical picture of concentric (onion-like) hypertrophy30. The prognosis of this event is poor. Before the introduction of ACE inhibitors (ACEI), almost all patients with severe renal insufficiency died within one year. Basic clinical symptoms of the renal crisis are: acute renal failure, without any warning signs, sudden
Therapy
• Discrete changes in urine sediment (mild proteinuria)

Diagnosis
Basic clinical findings in renal crisis are:
• Acute renal failure, without any preceding signs of significant renal disorder
• Sudden onset of moderate or severe hypertension, often accompanied by emergent hypertonic crisis, which may be as follows:
  o Hypertonic encephalopathy (headache, confusion, visual disorders)
  o Hypertonic retinopathy (hemorrhages, exudates)
  o Acute cerebrovascular event (CVE)
• Discrete changes in urine sediment (mild proteinuria and/or hematuria)

Therapy
Effective and rapid (within 72 h) control of blood pressure is the basic therapeutic measure. It provides stabilization of renal functions in almost 70% of patients and increase in one-year survival up to 80%. ACE inhibitors, which are contraindicated in a majority of other forms of acute renal failure, seem to be appropriate treatment of renal crisis in SSc. Captopril was the most frequently drug used in clinical trials, however, enalapril, ramipril and quinapril may provide similar protective effects. ACEI therapy may provide significant improvement of renal function to allow interruption in regular dialysis treatment and decrease the need of permanent dialysis.

The prognosis still remains unfavorable, mainly due to late diagnosis, and the rate of mortality is high.

Interstitial lung disease (ILD)
The most frequent interstitial changes are found in sub pleural areas of dorsal parts of both lower pulmonary lobes. In patients with moderate or severe restrictive disorder, interstitial changes propagate into upper parts of the lung field. Reduction in respiratory function, especially in forced vital capacity (FVC) can be detected early; reduction of FVC by 32% annually was detected in the first 2 years. ILD is associated with the high rate of mortality, which becomes conspicuous especially after five years of disease onset. Prognosis of patients with severe pulmonary impairment (FVC < 55% and DLCO < 40% of predicted normal values) is unfavorable; up to 42% die within 10 years of disease onset.

Diagnosis
Diagnosis is based on the chest X-ray, spirometry and high-resolution computed tomography (HRCT) findings evaluation, together with bronchoalveolar lavage or lung biopsy analyses.

Therapy: Lower doses of glucocorticoids and cyclophosphamide may be used in active alveolitis therapy.

Prognosis
The slowly progressing disease, which represents the majority of cases, has better prognosis than idiopathic interstitial pulmonary processes.

Pulmonary arterial hypertension (PAH)
It is the primary disorder of pulmonary arterioles. It accompanies systemic connective tissue diseases, mostly SSc, where it may appear together with ILD. With respect to severity, the disease may be classified as mild, moderate or severe. From the hemodynamic point of view, PAH is defined as precapillary pulmonary hypertension, with mean pulmonary artery pressure (MPAP) values ≥25 mmHg in rest. Gradually progressive exercise breathlessness and fatigue are the most frequent symptoms of PAH. Anginal pains reflect ischemia of the right heart ventricle, while syncope and presyncope are signs of low cardiac output. One interesting symptom, caused by dilated pulmonary artery stem related recurrent nerve palsy, is hoarseness. Advanced PAH is presented by the right heart failure, which may lead to the terminal right ventricular failure.

Diagnosis
Echocardiography is the key noninvasive diagnostic tool in the case of PAH suspicion. Currently, it is used as a routine screening procedure. By mean of echocardiography, we can evaluate size and function of the right ventricle and estimate the pressure in the pulmonary artery. Measurement of the tricuspid valve regurgitation gradient by Doppler echocardiography, and estimation of the right atrial pressure from the vena cava inferior size are fundamental for the pulmonary hypertension evaluation. A right heart catheterization should be indicated in symptomatic patients with suspected even mild pulmonary hypertension and it is crucial for PAH diagnosis confirmation. DETECT algorithm may be helpful in selection and indication of patients for this invasive procedure.

Therapy
Warfarin, as an anticoagulant therapy, is used, to prevent potential pulmonary embolism. Endothelin-1 receptor antagonists (e.g. bosentan, ambrisentan, macitentan), phosphodiesterase inhibitors (e.g. sildenafil, tadalafil) or intravenous prostanooids (epoprostenol) represent current specific treatment of PAH. Early initiation of therapy is advocated.

DERMATOMYOSITIS/POLYMYOSITIS (DM/PM)
It is a chronic autoimmune inflammatory disease affecting striated skeletal muscles (polymyositis) or muscles and skin, with elevation of respective muscle enzymes.
levels in serum. The disease may also afflict esophagus and lung, and rarely myocardium.

Predominant symptoms are proximal muscle weakness and typical skin finding in dermatomyositis heliotrope rash and Gottron's papules.

**Life-threatening events in DM/PM**

Pulmonary manifestations of the disease are major life threatening events in DM/PM (ref.39). They often appear in patients with positivity for anti-synthetase antibodies (e.g. anti-Jo). Breathlessness or respiratory insufficiency may appear due to the interstitial fibrosis (Fig. 1) or as a result of respiratory restriction caused by weakness of respiratory muscles. Patients may be endangered by an increased risk of aspiration and related aspiration pneumonia, which is listed as one of the leading causes of death in patients with restricted mobility and swallowing disorders40. Diffuse fibrosing alveolitis may switch to pulmonary fibrosis.

**Diagnosis**

Respiratory function test, radiography, HRCT and bronchoalveolar lavage are used as standard diagnostic tools together with the disease focused tests, like the examination of serum muscle enzyme levels, EMG, muscle biopsy and immunology.

**Therapy**

Therapy is based on high dose glucocorticoids, often in combination with immunosuppressant (azathioprine, methotrexate). High dose polyclonal immunoglobulins may be used as well, if indicated.

**Prognosis**

Pulmonary manifestation is one of the major negative prognostic factors in patients with PM/DM. For patients with DM, prognosis is worse than the disease related pulmonary involvement41.

**VASCULITIDES**

**Definition**

Vasculitides are a heterogeneous group of disorders, which are characterized by inflammation and necrosis of the vessel wall. The disease is often systemic, with multiorgan involvement. Primary vasculitis inflicts vessels of any size. Secondary vasculitis accompanies some rheumatic diseases, e.g. rheumatoid arthritis, SLE, or as a case of non-rheumatic disorders (sarcoidosis, malignancies, serious bacterial infections).

Signs and Symptoms: Pulmonary and renal afflictions are the most serious manifestations of vasculitis.

**Diagnosis**

Appropriate interpretation of multiple data from clinical, laboratory and imaging examinations is crucial for the disease diagnosis set-up. This can be reached through the close multidisciplinary cooperation of rheumatologist, pneumologist, nephrologist and clinical pathologist. Evaluation of antibodies against neutrophil cytoplasm (ANCA) is a valuable tool in so-called "ANCA-associated" vasculitis. Positivity of ANCA antibodies is typical for syndromes like granulomatosis with polyangiitis (with anti-proteinase-3 specific antibodies, cANCA) and other ANCA related vasculitis (with anti-myeloperoxidase specific antibodies, p-ANCA) (ref.42).

**Pulmonary impairment**

Serious pulmonary impairment appears mainly in ANCA associated vasculitis and pulmonary-renal (Goodpasture's) syndrome43.

**Granulomatosis with polyangiitis (formerly Wegener's granulomatosis)** is an ANCA positive vasculitis with primary involvement of small vessels and frequent respiratory tract impairment. Chronic rhinitis and sinusitis, associated with granular inflammation can be found in the upper respiratory tract. Breakdown of bone structure with the collapse of nasal tissue and the saddle nose formation may be the chronic sinusitis complication. Ulceration of the nasal mucosa or perforation of the nasal septum is frequent. Pulmonary impairment is reflected as fever, breathlessness, cough, hemoptysis, pleurodynia, and seldom by tracheal obstruction. Stenosis of the large airways may be the consequence of the repair processes, and can threaten the patient's life. Stridor is a typical sign of severe stenosis. Pulmonary infiltrates, sometimes extensive, are quintessential signs of the disease, which may be associated with the tissue decomposition process, pulmonary hemorrhage and respiratory failure that need artificial lung ventilation. Correlation between the disease exacerbation and infection of the lower respiratory tract has been well established44. Development of pulmonary fibrosis is a chronic sequel of the disease.

**Eosinophilic granulomatosis with polyangiitis** (Churg-Strauss syndrome) is necrotizing ANCA-related vasculitis, associated with asthma, eosinophilia and the granulo-
matous respiratory tract infection. The chest X-ray may detect fugitive pulmonary infiltrates. The histology examination shows predominant extravascular granulomas, rich in eosinophils, and signs of necrotizing vasculitis. Microscopic polyangiitis is the necrotizing ANCA positive vasculitis afflicting small vessels of the skin, kidneys and lung. Pulmonary hemorrhage is present in about 29% of patients (Fig. 2).

Pulmonary-renal (Goodpasture’s) syndrome is vasculitis with renal injury and alveolar hemorrhage. Positivity of the anti-GBM antibodies (antibodies against the glomerular basal membrane) is also detectable.

Renal impairment may be present, mainly in association with polyarteritis nodosa, ANCA-associated vasculitis, Henoch-Schönlein purpura and cryoglobulinemic vasculitis.

Polyarteritis nodosa was one of the first described vasculitis. It is a rather rare disease, which affects the arteries of small and medium gauge. In many cases, it is a sequella to viral B hepatitis. ANCA antibodies are positive in < 5% of cases. Renal injury is related to ischemic nephropathy, which may be accompanied by malignant hypertension. Renal failure may be the end-up outcome. Angiography often shows multiple vessel narrowing and microaneurysm of visceral and renal arteries. Renal insufficiency develops due to renal infarction and scarring of renal parenchyma. Aneurysm ruptures are infrequent. However, they can complicate renal biopsy outcome.

Renal ANCA positive vasculitis (granulomatosis with polyangiitis, eosinophil granulomatosis with polyangiitis, microscopic polyangiitis) affect the kidneys as a pauci-immune necrotizing glomerulonephritis with sickle cell formation. It can be clinically represented by rapidly progressing glomerulonephritis (RPGN). In contrast to lupus related nephritis, pauci-immune glomerulonephritis is characterized by the absence of striking deposits of immunoglobulins or complement particles.

Henoch-Schönlein purpura presents as rather frequent vasculitis of the small vessels with multiorgan impairment (skin, gastrointestinal tract, kidney, joints). It is ANCA negative, leucocytoclastic vasculitis. Microscopic or macrscopic hematuria with mild proteinuria, sometimes even nephrotic or nephritic syndrome are manifestations of renal impairment, with variable degree of renal insufficiency. Renal symptoms may appear earlier than the skin impairments. Histomorphology of the Henoch-Schönlein purpura related nephritis shows an IgA nephropathy with variable morphology, from a focal mesangial proliferative glomerulonephritis to necrotizing forms, with sickle cell formation. The characteristic features of the disease are the mesangial deposits of IgA.

Cryoglobulinemic vasculitis. Cryoglobulins are immunoglobulins, which reversibly precipitate under cool conditions (typically at 4 °C). The type I cryoglobulin presence is typical for the monoclonal gamapathies. It is composed of monoclonal immunoglobulin with no antibody activity. Cryoglobulins types II and III are characterized by the presence of mixed cryoglobulins, composed of at least 2 types of immunoglobulins. The type II cryoglobulins are monoclonal immunoglobulins with rheumatic activity against polyclonal IgG, the type III include polyclonal components. The type III mixed cryoglobulinemia is associated with systemic connective tissue disorders, leukemia, hepatobiliary disorders and infections. Close relationship between the mixed type II cryoglobulinemia and hepatitis C virus infection has been repeatedly confirmed. Skin purpura with leucocytoclastic vasculitis is a manifestation of mixed cryoglobulinemia. Skin lesions may be accompanied by large ulcerations or mutilations. There are other symptoms of the disease, like fatigue, fever, arthralgia and Raynaud’s syndrome, as well as peripheral neuropathy, accompanied by dysesthesia or anesthesia. Concomitant hepatosplenomegaly may often be a sign of hepatitis C. Proteinuria and microscopic hematuria are signs of renal impairment, frequently accompanied by renal insufficiency. As clinical symptoms, there may be nephritic syndrome, hypertension or nephrotic syndrome present. Severe renal insufficiency is more frequent in men and older patients. Laboratory exams confirm the presence of the mixed cryoglobulin, remarkable consumption of the complement components and positivity of rheumatoid factors. Renal biopsy is performed mostly in cases of type II mixed cryoglobulinemia, and shows membranoproliferative glomerulonephritis with relatively intense neutrophil and monocyte infiltrates and double contour of the basal membrane. In an acute phase
of the nephritic syndrome we can find intraluminal amorphous eosinophilic deposits too. Avoiding hypothermia is very important, especially in case of extracorporeal circulation use, e.g. hemodialysis, continuous elimination methods or ECMO.

**Therapy**

The vasculitis therapy differs according to individual diagnosis and extent of organ injury. Therapy includes glucocorticoids, cyclophosphamide, azathioprine, methotrexate, cyclosporine A, immunoglobulins, together with novel medication, such as mycophenolate mofetil and rituximab are. In cases of granulomatosis with polyangiitis and/or severe, life-threatening disorder, appropriate immunosuppression plays a key role, often in combination with plasmapheresis in cases of RPGN or pulmonary hemorrhage. Timely use of glucocorticoids, cyclophosphamide and plasmapheresis may significantly improve the patient’s prognosis.

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