Hypocomplementemic Urticarial Vasculitis Syndrome

An Interdisciplinary Challenge

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SUMMARY

Background: Chronic urticaria often points the way to the diagnosis of a systemic disease, particularly when urticarial vasculitis can be demonstrated. Hypocomplementemic urticarial vasculitis syndrome (HUVS) is considered to be an independent immunological disease.

Method: Selective literature review and consideration of the author’s own clinical experience.

Results and conclusions: The main manifestation of HUVS is chronic urticarial vasculitis with complement deficiency and the demonstration of C1q antibody in the serum. Multiple other organs are involved, sometimes severely. The diagnosis is confirmed by skin biopsy, which reveals leukocytoclastic vasculitis as a pathogenetic correlate of this systemic disease. Although HUVS is relatively rare, the medical specialists that might encounter it—ophthalmologists, rheumatologists, nephrologists, dermatologists, general practitioners, and pediatricians—should include it in their differential diagnoses whenever appropriate. Awareness of HUVS and rational diagnostic evaluation will lessen the chance of it being misdiagnosed as another type of systemic immunological disease and will reduce superfluous diagnostic testing in patients suffering from it.

Key words: urticaria, vasculitis, autoimmune disease, hypocomplementemia, chronic disease

Urticaria is a cutaneous vascular reaction with locally increased vascular permeability. It is clinically manifested by erythema and wheals and is accompanied by itching or a burning sensation. The wheals are usually transient, persisting for anything from a few minutes to a few hours. Urticarial vasculitis (UV) is a discrete disease entity, distinct from urticaria, in which the urticarial lesions persist for more than 24 hours and heal leaving brownish residues (1). On diascopy the cutaneous efflorescences generally display a central red spot, betraying the underlying vasculitis. Because leukocytoclastic vasculitis involves leakage of erythrocytes from blood vessels, hyperpigmentation remains after resolution of the urticarial lesion. In contrast, the exanthema seen in acute or chronic urticaria resolves completely within 24 hours (2).

Incidence

Some 2% to 20% of patients admitted to the hospital for investigation of chronic urticaria are found to have UV. Hypocomplementemic urticarial vasculitis syndrome (HUVS), however, a severe systemic form of urticarial vasculitis, is rare; its exact incidence is unknown. In one retrospective study at a single center, 18% of patients with biopsy-confirmed UV exhibited a hypocomplementemic form of the disease (3). HUVS affects women more frequently than men (ratio 2:1). The peak incidence is seen in the fifth decade of life, but HUVS has also been observed in children (4).

Case report

A 72-year-old woman had a 6-month history of night sweats, weight loss, worsening general health, urticarial skin lesions, and recurring conjunctivitis. Exhaustive dermatological, gastrointestinal, cardiological, hematono-cological, angiological, and radiological investigations failed to pinpoint the diagnosis. Attempts at treatment with antihistamines, triamcinolone lotions, and prednisolone (10 mg p.o.) were unsuccessful. Severe abdominal cramps and nausea led to hospital admission.
Physical examination revealed urticarial exanthema with areas of brownish hyperpigmentation on the trunk and extremities (Figure 1) accompanied by bilateral conjunctivitis. The patient was slightly sensitive to left lower abdominal and epigastric pressure and exhibited arthritis in the right ankle joint and first metacarpophalangeal joint that resolved spontaneously after 24 hours. Laboratory tests showed increased erythrocyte sedimentation rate (ESR) and c-reactive protein (CRP), anemia, an elevation of antinuclear antibodies to 1:1280—without demonstration of double-strand DNA antibodies, anti-extractable nuclear antigen (ENA), or antiphospholipid antibodies—but weakly positive mixed cryoglobulins with no monoclonal components, strongly decreased complement activity, and raised tumor marker CA125. Examination of the urinary sediment revealed isolated glomerular hematuria.

Gastroscopy, colonoscopy, gynecological examination, abdominal CT and echocardiography showed distinct ascites (granulocyte-rich exudate) and a discrete pericardial effusion with no indication of malignancy. The conjunctivitis was not improved by gentamicin eye drops. Histological examination of a repeat dermal biopsy sample from a fresh urticarial efflorescence on the thigh showed leukocytoclastic vasculitis with immunohistochemical demonstration of C1q deposits on the vascular endothelium. The material used for immune complex diagnosis was fixed in formalin and embedded in paraffin, with primary antibodies to IgA, IgG, IgM, C3c, and C1q. Antigen unmasking with proteases and a polymer-based detection system with horseradish peroxidase were employed to mark the binding sites of the primary antibodies brown. This procedure matches immunofluorescence for sensitivity and specificity. There were no bandlike deposits on the basal membrane. The chronic urticarial efflorescences with histological demonstration of small-vessel vasculitis and the strongly decreased complement factors pointed to the rare clinical picture of HUVS, particularly since all six minor criteria were fulfilled (Box 1).

Monotherapy with 2 g mycophenolate mofetil in short-term combination with 50 mg prednisolone swiftly resulted in complete remission for more than 24 months. Interestingly, the treatment for vasculitis also normalized the tumor marker level.

UV represents a disease continuum whose life-threatening end is occupied by HUVS. The UV spectrum embraces the following conditions:

- The mild form of disease with distinct urticarial exanthema and slight or absent systemic manifestations without hypocomplementemia (Figure 1a)
- Various special forms: AHA syndrome (21), Schnitzler syndrome (20), Cogan syndrome (19), Muckle–Wells syndrome (18) (Box 2)
- Life-threatening systemic vasculitis with hypocomplementemia and only mild urticarial skin lesions: HUVS (6).

HUVS is also known as McDuffie syndrome, after the researcher who published the first description (7).

**Box 1**

**Criteria for diagnosis of HUVS**

- **Major criteria**
  - Chronic urticarial exanthema
  - Hypocomplementemia

- **Minor criteria**
  - Leukocytoclastic vasculitis
  - Arthralgia and arthritis
  - Uveitis or episcleritis (or conjunctivitis)
  - Glomerulonephritis
  - Abdominal pain
  - Positive C1q antibody

*modified from (5); HUVS, hypocomplementemic urticarial vasculitis syndrome

**Figure 1**

a) Urticarial efflorescences on the upper body in hypocomplementemic urticarial vasculitis syndrome (HUVS)

b) Older efflorescences on the legs of the same patient: reddish hyperpigmentation as expression of urticarial vasculitis
Organ involvement

Hypocomplementemic urticarial vasculitis syndrome is a specific autoimmune disease with involvement of the skin, joints, kidneys, and gastrointestinal tract manifested by vasculitis and polyserositis. Occasionally other organ systems are involved.

Skin

In contrast to other forms of urticaria, the cutaneous efflorescences in UV are caused by leukocytoclastic vasculitis. The urticarial lesions differ from ordinary urticaria in that the wheals persist for more than 24 hours and leave brown residues after resolution (Table 1).

If the vasculitis also involves deeper vessels, angioedema may be found. Immunohistochemistry demonstrates immunoglobulin and complement deposits in the vessel walls or on the endothelium. Such deposits also occur together with vasculitis in systemic lupus erythematosus, but in lupus patients they are typically found along the basal membrane (the so-called lupus band) (4).

Joints

Arthralgia and arthritis of various joints are the most frequent systemic manifestations of HUVS. The joint pain wavers and is often no more than transient. The elbows, wrists, knees, and ankles are involved. Joint deformities are possible, in which case the cardiac valves are also affected (8).

Kidneys

Renal involvement is often only mild, but dialysis may be required (9). The frequently found proteinuria and hematuria with demonstration of acanthocytes (nephritic sediment) is seen histologically as membranous, membranoproliferative, or intra- and extracapillary glomerulonephritis. Nevertheless, renal biopsy is necessary only in proteinuria (>1 g/day) or in acute or chronic progressive renal failure. The renal involvement seems to be more severe in children (10, 11).

Lungs

Lung involvement goes along with shortness of breath, coughing, hemoptysis, pleural effusion, and chronic obstructive pulmonary disease (COPD) and is the most frequent cause of death among HUVS patients. The lungs are affected more severely in smokers (3).

Gastrointestinal tract

Around 30% of HUVS patients suffer gastrointestinal symptoms: pain, nausea, vomiting, diarrhea, sometimes also ascites in connection with serositis, hepatomegaly, and splenomegaly (12).
Other organs
Ocular involvement may be observed, with uveitis, episcleritis, and also conjunctivitis. The heart and the nervous system are occasionally affected (Table 2).

Pathophysiology
Although the pathophysiology of HUVS has not been fully clarified, the following cascade seems feasible: IgG antibodies to the collagen-similar regions of C1q form immune complexes and initiate the cascade. These C1q antibodies and immune complexes are also regularly detected in the plasma of patients with HUVS (13). The immune complexes activate the classical complement pathway into and around the blood vessels. The complement products generally lead to mast cell degranulation. The substances released thereby result in increased vascular permeability with urticaria and/or angioedema (14) and lead to histological demonstration of leukocytoclastic vasculitis in terms of a Coombs and Gell type III reaction.

Interestingly, anti-C1q autoantibodies have also been detected in pulmonary diseases. These antibodies can cross-react with surfactant and may thus be responsible for the lung involvement in HUVS (15, 16).

Diagnosis
Alongside chronic urticarial exanthema, the cornerstone of UV diagnosis is demonstration of leukocytoclastic vasculitis in dermal biopsy samples (17). This involves a leukocytoclastic reaction, vessel wall destruction, and deposits of fibrinogen. Immune complexes and complement should be visible in the blood vessels on immunohistochemical analysis. The more severe the findings, the more pronounced is the perivascular neutrophilic infiltrate (Figure 2).

For the diagnosis of HUVS, the typical cutaneous efflorescences must be accompanied by decreased complement levels. All HUVS patients exhibit extremely low levels of C1q. Through activation of the classical complement pathway, complement factors C3 and C4 are also nearly always markedly reduced. A C1q antibody can regularly be detected but is not specific for HUVS (13) (Box 3). The severity of the disease can be judged not just from the clinical signs and symptoms, but also from the degree of complement reduction.

Depending on what organs are affected, investigations such as abdominal sonography, echocardiography, chest x-ray, and lung function tests are required.

To detect any renal involvement, the urine should be examined by microscopy, protein quantification, and if necessary, protein analysis. If a nephritic syndrome is found, renal biopsy should be considered.

Definition
The criteria for diagnosis of HUVS were established by Schwartz (5) in 1982 (Box 1).

Angioedema, obstructive lung disease, or various neurological findings may also be present but are not listed among the minor criteria.

BOX 3

Typical laboratory findings in HUVS*
- ESR acceleration
- Hypocomplementemia with low C1q, C3, C4
- C1q antibodies
- ANA without anti-double-stranded DNA

* modified from (4); HUVS, hypocomplementemic urticarial vasculitis syndrome; ESR, erythrocyte sedimentation rate; ANA, antinuclear antibodies
Diagnostic criteria for systemic lupus erythematosus (SLE) by the American College of Rheumatology (ACR)*

- **Butterfly erythema**
  Fixed erythema, flat or elevated, in the buccal region, usually not involving the nasolabial folds

- **Discoid skin changes**
  Erythematous, raised skin lesions with adherent keratotic elements and follicular occlusion; atrophic scars can occur in old lesions

- **Photosensitivity**
  Skin reddening in the course of an unusual reaction to sunlight; occurrence reported by patients

- **Oral ulceration**
  Oral or nasal ulceration, usually painless; noted by a doctor

- **Arthritis**
  Non-erosive arthritis involving two or more peripheral joints, characterized by stiffness, swelling, and joint effusion

- **Serositis**
  a) Pleuritis—typical anamnesis for pleuralgia or pleural fremitus, detected by auscultation by a doctor, or demonstration of a pleural effusion, or
  b) Pericarditis—confirmed by ECG or by friction sounds, or by demonstration of a pericardial effusion

- **Renal disease**
  a) Persisting proteinuria of more than 0.5 g/day or greater than 3+, if no quantification is performed, or
  b) Cellular cylinders, erythrocyte, hemoglobin, granular, tubular, or mixed cylinders

- **Neural disease**
  a) Cramps—exclusion of drug induction or a metabolic disturbance, e.g., uremia, ketoacidosis, or electrolyte disturbances, or
  b) Psychosis—without obvious drug induction, exclusion of a metabolic disturbance, e.g., uremia, ketoacidosis, or electrolyte disturbances

- **Hematological disease**
  a) Hemolytic anemia with reticulocytosis, or
  b) Leukopenia, less than 4000 leukocytes/µL, demonstrated on two or more occasions, or
  c) Lymphopenia, less than 1500/µL on two or more occasions, or
  d) Thrombocytopenia, less than 100 000/µL without intake of potentially causative medication

- **Immunological disease**
  a) Anti-DNA: elevated titer of antibodies to native ds-DNA, or
  b) Anti-Sm: detection of antibodies to Sm antigens, or
  c) Demonstration of antiphospholipid antibodies:
    – elevated IgM or IgG anticardiolipin antibodies, or
    – demonstration of lupus anticoagulant by the standard test, or
    – false-positive serological test for syphilis, positive for more than 6 months, confirmed by means of a Treponema pallidum immobilization test or a fluorescence *Treponema* antibody absorption test

- **Antinuclear antibodies**
  Demonstration of an elevated antinuclear antibody titer on immunofluorescence or an equivalent test at a particular time, unconnected with any medication that may be associated with a so-called drug-induced lupus syndrome

* If four or more criteria are fulfilled, systemic lupus erythematosus is probable.
Modified from [62]
UV has taken a severe course, with multiple organ exclusion and hypocomplementemia is present, then the syndrome of AHA syndrome.

Concentration or function. Little is known about the causes of C1 esterase inhibitor displays no abnormality of concentration or function.

As a special form of chronic urticaria, diagnosis of UV requires dermal biopsy with demonstration of leukocytoclastic vasculitis and a perivascular neutrophilic infiltrate. If UV is present, one must establish whether it is isolated or exists in the context of a syndrome.

In Muckle–Wells syndrome there is a mutation of the NLRP3 gene on chromosome 1q44, which codes the protein cryopyrin. Under physiological conditions this protein is activated in the presence of danger and leads to release of interleukin-1. Possibly the mutated protein cannot be inactivated quickly enough, resulting in excessive reactions.

Clinically, besides the UV, once every few weeks the patients display a 12- to 36-hour-long phase of intermittent fever with painful joints, progressive hearing loss, and secondary amyloid A protein amyloidosis with renal involvement (18).

Cogan syndrome preferentially affects young adults in the third decade of life. Following a rheovirus-III infection, antibodies to the virus core protein can cross-react with the inner ear and eye. The clinical presentation resembles that of Menière disease, with fainting attacks, hearing loss, and interstitial keratitis with reddened, painful eyes and visual impairment (19).

In Schnitzler syndrome UV is combined with a monoclonal IgM gammopathy. Apart from intermittent attacks of fever, the clinical manifestation are bone and joint pains and hyperostoses. A lymphoproliferative disease cannot be detected. The pathogenesis and the part played by IgM are unknown (20).

In arthritis, hives, and angioedema (AHA) syndrome, UV is accompanied by angioedema (21). The C1 esterase inhibitor displays no abnormality of concentration or function. Little is known about the causes of AHA syndrome.

If the above-mentioned syndromes have been excluded and hypocomplementemia is present, then the UV has taken a severe course, with multiple organ involvement, and the differential diagnoses are systemic lupus erythematosus (SLE) and the mixed cryoglobulinemias (22). The active forms of all three diseases feature low complement levels and involvement of multiple organs.

Differentiation from SLE is sometimes difficult, because 50% of patients with HUVS show positive antinuclear antibodies and often also fulfill the classification criteria of the American College of Rheumatology (Box 4).

The patient whose case is presented above, however, fulfilled only three of the 11 SLE criteria. Inversely, in 50% of patients with confirmed UV the cause is SLE (3). The C1q antibodies are also not specific for HUVS and may be found in one third of patients with SLE (13).

In addition to the ACR criteria, HUVS and SLE can often be differentiated on the basis of double-stranded DNA antibodies and the typical clinical presentations. High titers of double-stranded DNA antibodies or the demonstration of SLE-typical ENA speak against HUVS.

HUVS can be distinguished from Sharp syndrome, another possible differential diagnosis amongst the collagenoses, by demonstration of the U1-RNP antibody.

The differentiation between HUVS and mixed cryoglobulinemia type III also exhibits overlaps, in that cryoglobulins may occur as an epiphenomenon in both SLE and HUVS (22). Differentiation from mixed cryoglobulinemia type II, however, is possible by the demonstration of a monoclonal immunoglobulin in the cryoprecipitate and on the basis of the usually higher cryoglobulin concentration.

**Treatment**

The appropriate treatment of UV and HUVS is determined by the severity of the disease. Because the disease spectrum reaches all the way from simple UV with skin lesions without hypocomplementemia to HUVS with severe systemic involvement, the treatment options are similarly diverse. The rarity of the disease means there is little evidence, and the treatment recommendations are based exclusively on reports of single cases or small series.

Antihistamines represent the cornerstone of treatment for patients who have UV with only cutaneous lesions, serving to control the itching. Antihistamines seldom suffice alone, however, because they do not bring the immune complex formation under control, intervene very late in the inflammation cascade, and cannot alter the course of the disease (23).

Glucocorticoids (GC) are the agents most frequently employed to combat the inflammation and immune complex formation. The dosage depends on the severity of the disease. Patients with hypocomplementemia initially almost always require GC. Because of the well-known adverse effects numerous other immunosuppressives are combined with GC, including methotrexate, azathioprine, cyclophosphamide, cyclosporin, and mycophenolate mofetil. In agreement with the observations of investigators from Berlin (24), this...
last-mentioned drug was notably successful in the patient whose case is presented above.

In highly active disease plasmapheresis can also be employed. This reliably reduces the circulating immune complexes, but the effect is only transitory and the symptoms re-emerge a few days later unless C1q antibody production is blocked by means of immunosuppressives.

Dapsone has also been used in the treatment of HUVS. Because of the wide range of disease severity, however, the role of dapsone has not been clearly delineated. It may be possible to employ dapsone together with GC and an immunosuppressive in a three-pronged approach (25). Newly developed treatment methods have been successfully employed in the special forms of UV. For instance, there have been repeated reports of the successful use of the interleukin-1-receptor antagonist anakinra in Schnitzler syndrome (20) and Muckle–Wells syndrome (18). In contrast, the B-cell antibody rituximab yielded no therapeutic benefit (20).

**Conflict of interest statement**
The authors declare that no conflict of interest exists according to the guidelines of the International Committee of Medical Journal Editors.

Manuscript received on 28 January 2008, revised version accepted on 21 April 2009.

Translated from the original German by David Roseveare.

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