The Diagnosis and Treatment of Systemic Lupus Erythematosus

Annegret Kuhn, Gisela Bonsmann, Hans-Joachim Anders, Peter Herzer, Klaus Tenbrock, Matthias Schneider

SUMMARY

Background: Systemic lupus erythematosus (SLE) is an autoimmune disease with a prevalence of 36.7/100 000 in Germany and a female/male ratio of 4:1. The clinical course is variable, with a broad spectrum of organ manifestations; lupus nephritis develops in about half of all patients.

Methods: This review is based on a selective search of PubMed and the Cochrane Library, including current guidelines and expert recommendations.

Results: Assessment of clinical symptoms, laboratory findings, and optional biopsy results are the basis for early diagnosis of SLE. All patients should be treated with antimalarials as soon as the diagnosis is confirmed. In particular, hydroxychloroquine is associated with a higher rate of remission, fewer relapses, and reduced damage in the course of the disease, even in lupus nephritis. High-dose glucocorticoids should be given only when acutely indicated; immunosuppressives such as azathioprine, methotrexate, or mycophenolate mofetil may be administered to reduce glucocorticoids, according to the EULAR recommendations. Belimumab was recently approved as add-on therapy in autoantibody-positive SLE patients with high disease activity unresponsive to standard treatment. Short-term induction pulse therapy with low-dose intravenous cyclophosphamide, as well as continued mycophenolate mofetil treatment are advances in lupus nephritis.

Conclusion: The long-term prognosis for SLE has improved markedly in recent decades because of earlier diagnosis and optimized treatment. Further research and randomized controlled trials are needed for the development of specifically targeted therapies.

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Systemic lupus erythematosus (SLE) is a heterogeneous autoimmune disease that may involve many different organs and display a variable clinical course. The diagnosis of SLE is based on characteristic clinical findings of the skin, joints, kidneys, and the central nervous system, as well as on serological parameters such as antinuclear antibodies (A1 A), in particular antibodies to dsDNA (e1). The various clinical symptoms do not always occur simultaneously and may develop at any stage of the disease. In the early stages, physicians from various disciplines often propose several differential diagnoses, or identify only one aspect of the disease without recognizing the symptoms as part of SLE (1, e2). Fever, fatigue, and arthralgia are the most frequently occurring non-specific symptoms at disease onset; additional joint swelling or a “butterfly rash”—particularly in women of childbearing age—should prompt consideration of SLE (2). The aim of this article is to provide an updated review on the diagnosis and treatment of SLE, based on a selective survey of the literature in PubMed and the Cochrane Library, including current guidelines and the recommendations of experts with extensive experience in the management of this disease.

Epidemiology and Prognosis

The prevalence of SLE in Germany in the year 2002 was 36.7/100 000, with a 4:1 ratio of women to men (3). The prevalence of pediatric-onset SLE is probably lower by a factor of ten (e3). The disease often begins in puberty; if SLE is diagnosed in patients under the age of 5 years, a rare monogenic form may be present. The survival rate has risen significantly in recent decades (1955 vs. 2003: 5-year survival rate 5% vs. 95%; 10-year survival rate 0% vs. 92%), mainly due to earlier diagnosis and improved management (4, 5, e4). During the first years after the onset of SLE, mortality is increased mainly due to disease activity and bacterial infection as a result of high glucocorticoid dosage (e5, e6), while cardiovascular complications predominate in the period beginning 5 years after initial diagnosis (e7, e8).

Classification criteria

The criteria of the American College of Rheumatology (ACR), first published in 1982 and revised in 1997, can be applied for the classification of SLE (6, 7, e9). Four
of the 11 criteria have to be fulfilled for a diagnosis of SLE. As 4 of the criteria include mucocutaneous lesions, the application of the ACR criteria without analysis of autoantibodies may result in an overestimation of SLE (8, e2). Therefore, the Systemic Lupus International Collaborating Clinics (SLICC) group developed a new set of classification criteria in 2012 (Box 1) (9). Currently both sets of criteria (ACR and SLICC) are often applied simultaneously.

**Diagnosis**

**Clinical manifestations**

Cutaneous manifestations occur in about 75% of patients with SLE in the course of the disease, and are the first sign in a quarter of cases (e10). Based on clinical and histological criteria, the skin lesions are divided into lupus erythematosus (LE)-specific and LE-non-specific manifestations (e11, e12). The most frequent LE-specific manifestation is the acute cutaneous lupus erythematosus (ACLE), which may occur as a butterfly rash or in the form of a generalized maculopapular exanthema. Discoid lupus erythematosus (DLE) displays a chronic course, typically characterized by inflammatory erythematous plaques with follicular hyperkeratosis and scarring (Figure 1a), leading to irreversible alopecia in hairy areas (Figure 1b). Approximately 5% of patients with DLE who initially show no systemic organ involvement will develop SLE. Subacute cutaneous lupus erythematosus (SCLE) is characterized by symmetric, annular, polycyclic, and/or papulosquamous/psoriasiform skin lesions without scarring on sun-exposed areas of the back, chest, and extensor surfaces of the arms (Figure 1c). The LE-non-specific manifestations include, among others, vascular skin lesions (e.g., periungual telangiectasia, livedo racemosa, Raynaud syndrome).

The musculoskeletal system is involved in around 90% of patients with SLE. In addition to myalgia and arthralgia, arthritis of small and large joints may occur (2). Tendovaginitis and synovitis can be detected by sonography and/or magnetic resonance imaging. Chronic recurring arthritis—particularly of the finger joints—may lead to joint deformation without radiological signs of erosions. This deforming but non-destructive joint disease is termed Jaccoud arthropathy (Figure 1d). About 50% of patients with SLE develop renal involvement (10). Lupus nephritis is a glomerular nephritis, typically with proteinuria and erythrocyturia (particularly dysmorphic erythrocytes) as well as erythrocyte cylinders in the urinary sediment (Figure 2a). Proteinuria should be quantified in a 24-hour urine sample; a finding of >0.5 g/day is usually an indication for renal biopsy. The 2003 revision of the histological classification of lupus nephritis distinguishes six forms (class I to VI), which require a differentiated treatment strategy (7, 11). The most commonly occurring cardiovascular manifestations of SLE include Libman–Sacks endocarditis, as well as pericarditis or myocarditis and coronary arteritis. Early occurrence of arteriosclerosis and the associated complications are

<table>
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**Classification of SLE: the Systemic Lupus International Collaborating Clinics (SLICC) Classification Criteria***

- **Clinical criteria**
  - Acute cutaneous lupus erythematosus (including "butterfly rash")
  - Chronic cutaneous lupus erythematosus (e.g., localized or generalized discoid lupus erythematosus)
  - Oral ulcers (on palate and/or nose)
  - Non-scarring alopecia
  - Synovitis (≥ 2 joints) or tenderness on palpation (≥ 2 joints) and morning stiffness (≥ 30 min)
  - Serositis (pleurisy or pericardial pain for more than 1 day)
  - Renal involvement (single urine: protein/creatinine ratio or 24-hour urine protein, >0.5 g)
  - Neurological involvement (e.g., seizures, psychosis, myelitis)
  - Hemolytic anemia
  - Leukopenia (<4000/μL) or lymphopenia (<1000/μL)
  - Thrombocytopenia (<100 000/μL)

- **Immunological criteria**
  - ANA level above laboratory reference range
  - Anti-dsDNA antibodies
  - Anti-Sm antibodies
  - Antiphospholipid antibodies (anticardiolipin and anti-β2-glycoprotein I [IgA-, IgG- or IgM-] antibodies; false-positive VDRL [Venereal Disease Research Laboratory] test)
  - Low complement (C3, C4, or CH50)
  - Direct Coombs test (in the absence of hemolytic anemia)

* Modified (short form) after (9).

For classification as SLE, four criteria (at least one of them clinical and at least one immunological) have to be fulfilled or lupus nephritis has to be diagnosed histologically in the presence of ANA or anti-dsDNA antibodies. The SLICC criteria are not diagnostic criteria. SLE, systemic lupus erythematosus.

* Modified (short form) after (9).
the major causes of mortality in SLE (4). The most common form of pulmonary involvement is dry or wet pleurisy (40 to 60%); lupus pneumonitis and pulmonary fibrosis are rare (2). The central nervous system is affected in about 15 to 50% of patients with SLE, but due to the low specificity (e.g., headache) and high variability of the symptoms, their identification as part of SLE often proves difficult (2, 12). Besides neuropsychiatric manifestations such as cognitive deficits, psychosis, and depression, organic brain syndromes, epileptic seizures, and transverse myelitis may occur, as well as vascular events (e.g., vasculitis, arterial/venous thrombosis in the case of antiphospholipid antibodies) and peripheral neuropathy (12, e13).

Laboratory diagnostics
If SLE is suspected based on clinical findings, laboratory testing can support the diagnosis. At first, a screening laboratory test is recommended (2, 8, 10, 13) (Box 2). A high erythrocyte sedimentation rate is characteristic for active SLE; C-reactive protein is usually normal or only slightly elevated. A standard or differential blood count may reveal cytopenias such as thrombocytopenia and/or leukopenia and lymphopenia, as well as further hematological changes such as autoimmune hemolytic anemia (e13). Renal parameters should include serum creatinine, urinary status and sediment. Antinuclear antibodies (AN A) should be analyzed by indirect immunofluorescence tests (HEp-2 cells) (Figure 2b). In patients with positive AN A and a homogeneous fluorescence pattern it is useful to determine anti-dsD| A antibodies. The findings of ELISA (high sensitivity, low specificity) should be confirmed by radioimmunoassay (RIA, also known as Farr assay; high specificity, suitable for monitoring) (14, e14). An alternative to the RIA, which is not available in all laboratories due to its requirement of radioactive material,
is the *Crithidia luciliae* immunofluorescence test (high specificity, low sensitivity). In addition to anti-dsDNA antibodies (prevalence 70 to 98%), anti-Sm antibodies (prevalence 14 to 40%) are specific marker antibodies of SLE (14, 15, e10). C3 and C4 should be determined as indicators of complement consumption or deficiency. The course of SLE is characterized by exacerbations and remissions; however, the AI A titer does not correlate with disease activity. In contrast, anti-dsDNA antibodies often rise several months before a disease flare, in parallel with a decrease of complement factors (16, 17). As a consequence, disease activity should be closely monitored—particularly with regard to renal involvement.

**Additional diagnostics**

Further diagnostic investigations depend on the symptoms of SLE and should be carried out in cooperation with medical specialists from the appropriate disciplines. In the case of cutaneous manifestations, a dermatologist should be consulted for a biopsy of the skin and/or oral mucous membrane lesions (histology/immunofluorescence) for purposes of differential diagnosis (e15). For further investigations see *Box 3*.

**Assessment of activity and damage**

For the determination of disease activity in SLE, various scoring systems have been developed (SLEDAI, SLAM, BILAG, ECLAM) that mainly include clinical findings and laboratory parameters (18). These scores enable standardized evaluation of disease activity, which is the basis for every therapy decision. Therefore, the European League Against Rheumatism (EULAR) recommends the use of a standardized score for assessment of disease activity at every visit. It is important to distinguish activity and damage, as the damage cannot be treated with immunosuppressive agents. The damage should be assessed annually using the SLICC/ACR Damage Index for SLE (Systemic Lupus International Collaborating Clinics/American College of Rheumatology) (19). The CLASI (Cutaneous Lupus Erythematosus Disease Area and Severity Index) and the RCLASI (Revised Cutaneous Lupus Erythematosus Disease Area and Severity Index) were developed and validated to assess disease activity and damage of mucocutaneous manifestations (20, e16).

**Treatment**

The treatment of SLE is based on the results of only few randomized controlled trials. Furthermore, the inclusion criteria of these studies were not uniform, with heterogeneous disease severity and low patient numbers. Nevertheless, internationally consented recommendations for the treatment of patients with SLE have been developed (*Tables 1 and 2*) (10, 12, 21).

**Antimalarial agents**

In every patient with SLE treatment with antimalarials is recommended unless there are contraindications (21–23). The action of antimalarials is based on, among other factors, the inhibition of activation of intracellular toll-like receptors (e17). Hydroxychloroquine and chloroquine are licensed for the treatment of SLE. Apart from their good efficacy against arthritis and LE-specific skin lesions (8), antimalarials maintain SLE in remission, are associated with fewer disease flares, and reduce damage in the course of the disease (23, e18). In lupus nephritis, cohort studies have demonstrated positive effects for hydroxychloroquine or chloroquine administered before (24), and for hydroxychloroquine as adjunct to standard treatment after the diagnosis of renal involvement (10, 25, e19). Furthermore, the positive impact of antimalarials on lipid and glucose metabolism have been described, as well as a reduction of thromboembolisms with a favorable influence on cardiovascular risk in SLE, and antineoplastic effects have been discussed (22, 23). Based on the experience of experts, the continuation of hydroxychloroquine treatment during pregnancy is recommended,
### BOX 2

**Investigations in suspected systemic lupus erythematosus (SLE) and monitoring after diagnosis**

- **Screening laboratory tests**
  - Erythrocyte sedimentation rate
  - Blood count, differential blood count
  - Creatinine
  - Urinary status and sediment
  - Antinuclear antibodies (ANA) (HEp-2 cell test with fluorescence pattern)

- **Further laboratory tests after positive screening** (particularly in case of positive ANA)
  - Further differentiation of ANA (particularly anti-Sm, -Ro/SSA, -La/SSB, -U1RNP antibodies, etc.)
  - Anti-dsDNA antibodies (ELISA; confirmation by radioimmunoassay or immunofluorescence test with *Crithidia luciliae*).
  - Complement C3, C4
  - Antiphospholipid antibodies, lupus anticoagulant
  - Glomerular filtration rate; 24-hour urine (if urine protein positive), alternatively: protein/creatinine ratio in single urine sample; investigation for dysmorphic erythrocytes in sediment
  - Liver enzymes: lactate dehydrogenase; creatine kinase in presence of muscular symptoms
  - Further laboratory tests depending on clinical symptoms
  - Screening for comorbidities
  - Assessment of vaccination status

- **Follow-up** (SLE: every 3 to 6 months depending on disease course; lupus nephritis: initially every 2 to 4 weeks for the first 2 to 4 months)
  - Medical history (including new symptoms, comedication, infections), physical examination
  - Evaluate disease activity with standardized score
  - Evaluate damage according to standardized score (1 ×/year)
  - Repeat screening for comorbidities (at least 1 ×/year)
  - Ocular examination in patients taking hydroxychloroquine or chloroquine: baseline, then every 6 months (currently being revised by the German Society of Rheumatology in light of recommendations from the USA) (e30, e31)
  - Laboratory tests
    - Erythrocyte sedimentation rate
    - C-reactive protein (in suspected infection or pleurisy)
    - Blood count, differential blood count
    - Creatinine
    - Liver enzymes
    - Urinary status (protein/creatinine ratio, 24-hour urine and microscopic examination of urinary sediment as needed)
    - Complement C3, C4
    - Anti-dsDNA antibodies
    - Instrument-based diagnostics as needed

*1 Modified after (2, 8), *2 modified after (2, 8, 10, 13, 21)

### BOX 3

**Organ-specific diagnostics as required**

- **Skin/oral mucous membrane**
  - Biopsy: histology, immunofluorescence if indicated

- **Joints**
  - Conventional X-ray
  - Arthrosonography
  - Magnetic resonance imaging (MRI)

- **Muscle**
  - Creatine kinase
  - Electromyography
  - MRI
  - Muscle biopsy

- **Kidney**
  - Sonography
  - Renal biopsy

- **Lung/heart**
  - Chest X-ray
  - Thoracic high-resolution computed tomography (HR-CT)
  - Lung function test including diffusion capacity
  - Bronchoalveolar lavage
  - (Transesophageal) echocardiography
  - Cardiac catheterization
  - Cardiac MRI
  - Myocardial scintigraphy
  - Coronary angiography

- **Eye**
  - Funduscopy/special investigations in patients on antimalarials

- **Central and peripheral nervous system**
  - Electroencephalography
  - Primarily cranial MRI, special MRI techniques if indicated
  - Computed tomography
  - Cerebrospinal fluid analysis
  - Transcranial Doppler/angiography
  - Neuropsychiatric examination
  - Measurement of nerve conduction velocity
as SLE patients on hydroxychloroquine show lower disease activity and fewer exacerbations and need lower doses of glucocorticoids at the time of birth (26, e20). Hydroxychloroquine can also be continued during breastfeeding (26, e21).

The dose of hydroxychloroquine and chloroquine depends on the maximal daily dose, calculated on the basis of ideal body weight (Table 1). If the actual body weight is lower than the ideal body weight, actual body weight should be used for calculation of the maximum daily dose (27). The dreaded side effect of irreversible retinopathy (bull’s eye maculopathy) was, particularly in earlier years, associated with excessive daily dosages (28, e22). In advanced renal failure (GFR <30 mL/min), the daily dose of antimalarial medication should be modified accordingly; in patients on dialysis it should be reduced to 15% of the standard dose (10, 22).

Hydroxychloroquine is preferentially used in SLE, but in case of intolerance or lack of efficacy chloroquine can be administered instead (8). The optimal efficacy of antimalarials is often not observed before 3 to 6 months of therapy. LE-specific skin lesions, however, may respond after 4 to 6 weeks (8, e22).

If no improvement of the cutaneous manifestations is seen after 3 to 6 months of treatment, hydroxychloroquine or chloroquine can be combined with the antimalarial agent mepacrine (synonym quinacrine; off label, no ocular toxicity) in a dosage of no more than 100 mg/day (8, e22).

Smoking has been suggested to decrease the effect of antimalarials; therefore, SLE patients should be advised to cease smoking completely. A few case reports and our own experience even show that therapy-refractory skin lesions were thereby resolved (e23, e24).

**Topical treatment**

Glucocorticoids are the topical treatment of choice for skin lesions in SLE (evidence level III) (8). Class IV glucocorticoids (e.g., clobetasol) can be applied to the scalp, palms, and soles, whereas in other areas only class II (e.g., methylprednisolone aceponate) and class III (e.g., mometasone furoate) glucocorticoids are recommended. Due to the adverse effects (e.g., atrophy, telangiectasia, perioral dermatitis) glucocorticoids should be administered only intermittently and not long term, particularly not for butterfly rash.

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**TABLE 1**

<table>
<thead>
<tr>
<th>Indication</th>
<th>Medication</th>
<th>Level of evidence</th>
<th>Strength of statement</th>
<th>Dosage</th>
</tr>
</thead>
<tbody>
<tr>
<td>First line and basic treatment</td>
<td>Hydroxychloroquine or Chloroquine</td>
<td>2 (21)</td>
<td>A (21)</td>
<td>≤ 6.0–6.5 mg/kg ideal body weight/day</td>
</tr>
<tr>
<td></td>
<td>If indicated, initial non-steroidal anti-inflammatory drugs and/or glucocorticoids</td>
<td>–</td>
<td>D</td>
<td>≤ 3.5–4.0 mg/kg ideal body weight/day</td>
</tr>
<tr>
<td></td>
<td>Azathioprine or methotrexate or mycophenolate mofetil*</td>
<td>4 (21)</td>
<td>B (21)</td>
<td>2–3 mg/kg body weight/day</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2 (21)</td>
<td>A (21)</td>
<td>15–20 mg/week (preferably s.c.)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>6 (21)</td>
<td>D (21)</td>
<td>2 g/day</td>
</tr>
<tr>
<td>If no response or no reduction of glucocorticoids ≤ 7.5 mg possible in the long term</td>
<td>Belimumab</td>
<td>–</td>
<td>B (21)</td>
<td>10 mg/kg body weight i.v. infusion (1 h) initially, then after 14 days and subsequently every 4 weeks</td>
</tr>
</tbody>
</table>

Remarks:

– According to expert opinion, not only low-dose prednisone but also hydroxychloroquine and azathioprine (particularly in lupus nephritis [10]) can be administered in pregnancy (e20).
– In case of comedication with mycophenolate mofetil and proton pump inhibitors, the bioavailability of mycophenolate mofetil is reduced; a switch to mycophenolic acid is advisable (e32).
– Proton pump inhibitors may lower the efficacy of hydroxychloroquine/chloroquine (e33).
– Treatment and monitoring instructions of the DGRh (in German) for the above-mentioned medications for use by patients and physicians can be found at www.dgrh.de/therapieueberwachen.html

DGRh, German Society of Rheumatology
* In patients with lupus nephritis, mycophenolate mofetil and i.v. cyclophosphamide have recently shown comparable efficacy with regard to total non-renal disease activity (39, 40)
An alternative is the off-label use of topical calcineurin inhibitors (tacrolimus ointment [level of evidence I], pimecrolimus cream [level of evidence II]). These agents can be applied as long-term treatment without the above-mentioned risks associated with the extended use of glucocorticoids (e.g. in butterfly rash, lupus erythematosus tumidus, initial DLE) (29, e25).

**Immunosuppressive treatment**

In patients without organ-threatening manifestations (e.g., LE-specific skin lesions, arthritis, pleurisy), long-term treatment with antimalarials should be sufficient. Due to the delayed onset of action of antimalarials, most patients temporarily need additional, short-term effective medication, usually non-steroidal anti-inflammatory drugs or glucocorticoids. If glucocorticoids cannot be reduced to a dosage of 5 to 7.5 mg/day prednisolone equivalent or discontinued within a reasonable period of time, the EULAR recommends—even for patients without organ involvement—administration of immunosuppressives such as azathioprine, methotrexate, or mycophenolate mofetil to spare glucocorticoids (21, 30). Methotrexate may have a favorable effect on joint and skin lesions and on general disease activity (8, 21, 31).

### TABLE 2

**Treatment recommendations on the example of proliferative lupus nephritis class III–IV in systemic lupus erythematosus with active organ involvement (ISN/RPS, International Society of Nephrology/Renal Pathology Society 2003 Classification of Lupus Nephritis) (11)**

<table>
<thead>
<tr>
<th>Indication</th>
<th>Medication</th>
<th>Level of evidence</th>
<th>Strength of statement</th>
<th>Dosage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Continuation of treatment with antimalarials</td>
<td>Hydroxychloroquine</td>
<td>1 (10)</td>
<td>C (10)</td>
<td>≤ 6.0–6.5 mg/kg ideal body weight/day</td>
</tr>
<tr>
<td>Induction therapy</td>
<td>Combination of glucocorticoids</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>with: mycophenolate mofetil</td>
<td>1 (10)</td>
<td>A (10)</td>
<td>Begin with methylprednisolone 500–750 mg i.v. on 3 consecutive days (level of evidence 3, strength of statement C); then glucocorticoids per os 0.5 mg/kg body weight/day for 4 weeks with subsequent tapering (strength of statement C)</td>
</tr>
<tr>
<td></td>
<td>or cyclophosphamide (low-dose) i.v.</td>
<td>1 (10)</td>
<td>B (10)</td>
<td>3 g/day or mycophenolic acid 2.16 g/day for 6 months, respectively (especially with marked proteinuria)</td>
</tr>
<tr>
<td></td>
<td>or azathioprine</td>
<td>2 (10)</td>
<td>B (10)</td>
<td>Total dose of 3 g (6 × 500 mg every 2 weeks) over 3 months</td>
</tr>
<tr>
<td>Maintenance treatment after response to induction therapy</td>
<td>Combination of low-dose glucocorticoids with: mycophenolate mofetil</td>
<td>1 (10)</td>
<td>A (10)</td>
<td>2 mg/kg body weight/day, in selected patients without adverse prognostic factors or if mycophenolate mofetil or cyclophosphamide are contraindicated, not tolerated, or unavailable</td>
</tr>
<tr>
<td></td>
<td>or azathioprine*</td>
<td>1 (10)</td>
<td>A (10)</td>
<td></td>
</tr>
<tr>
<td>Refractory to therapy, or contraindication</td>
<td>Calcineurin inhibitors (cyclosporine A, tacrolimus)</td>
<td>–</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Rituximab (anti-CD20)</td>
<td>–</td>
<td>–</td>
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</tr>
</tbody>
</table>

**Remarks:**
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DGRh, German Society of Rheumatology

effective option in SLE (e26). In 2012, belimumab was approved as an adjunct therapy for adult patients with autoantibody-positive SLE who despite standard therapy show high disease activity, intolerance of other treatments for SLE, or an unacceptably high need for glucocorticoids (Table 1) (e27). The most frequently occurring adverse effects include nausea, diarrhea, and bacterial and viral infections (e.g., bronchitis, cystitis, and pharyngitis), as well as hypersensitivity/infusion reactions. Data on the efficacy of belimumab in routine clinical practice are limited (32, e26).

The next level of treatment is indicated in the presence of active organ involvement, particularly when the kidneys and/or central nervous system are affected. The EULAR has recently published guidelines for the management of neuropsychiatric manifestations of SLE (12). The main goal is to identify cases of antiphospholipid syndrome, which requires anticoagulation and not immunosuppression. Otherwise, immunosuppressive treatment of SLE with organ involvement is mainly based on the experience with lupus nephritis (10, 33).

Mycophenolate mofetil (or mycophenolic acid in an equivalent dose) and low-dose intravenous cyclophosphamide in combination with glucocorticoids are available as induction therapy for proliferative lupus nephritis (class III–IV), depending on the best risk–benefit ratio (Table 2) (10). For patients who respond to initial treatment, the recommended maintenance therapy is lower immunosuppression, with either mycophenolate mofetil/mycophenolic acid or azathioprine for at least 3 years in combination with low-dose prednisone. Thereafter, a gradual reduction of the medication can be attempted, beginning with tapering of glucocorticoids. Until recently, mycophenolate mofetil or mycophenolic acid were described as superior to azathioprine for the maintenance therapy of lupus nephritis, but the 10-year data from the MAIİ TAIİ study and current meta-analyses do not confirm the previous findings (Table 2) (34).

Although SLE is usually more severe in childhood and adolescence, there are no randomized controlled trials of treatment in this age group. The treatment is based on data from adults and begins with hydroxychloroquine and glucocorticoids (35). In the presence of serious organ involvement (kidneys, central nervous system), early pulse therapy with cyclophosphamide and/or glucocorticoids is indicated. Mycophenolate mofetil is increasingly becoming an alternative to cyclophosphamide. Azathioprine and cyclosporine can be used as steroid-sparing medications. Patients are currently being recruited for a pediatric study of treatment with belimumab (e28).

**Adjunct treatment**

Almost as important as immunosuppression is the co-medication, which is determined by the comorbidities (infections, arteriosclerosis, hypertension, dyslipidemia, diabetes, osteoporosis, avascular necrosis, and malignancies, among others) and the damage that has already occurred (13, 21, 30). Weight control and adequate exercise are further important aspects. Depending on the medication prescribed and the clinical situation, the use of low-dose aspirin, calcium, vitamin D, bisphosphonates, statins, and blood pressure drugs (especially ACE inhibitors in proteinuria) should be considered. Immunosuppressed patients should be vaccinated (particularly against influenza and pneumococci); vaccinations with live attenuated vaccines are contraindicated (Box 2) (36).

Prophylactic measures comprise ultraviolet (UV) light protection, abstinence from smoking, and avoidance of medications known to trigger SLE (e.g., hydralazine and Tİ F-alpha antagonists; the reader is referred to the literature) (8, 37). Consistent light protection includes wearing sun protective clothing, keeping the head covered, and using sun blockers with chemical and/or physical UV-A/UV-B filters (SPF 50+). These should be applied 20 to 30 min before UV exposure in sufficient amounts (2 mg/cm²) (38, e29). SLE patients should be informed that UV-A radiation penetrates window glass (including that in vehicles).

**Conclusion**

Advances in diagnostics, optimized treatment (also of comorbidities), and regular monitoring of disease activity and damage have clearly improved the prognosis of SLE. The goal of treatment is remission or at least minimization of disease activity and prevention of flares. Nowadays, antimalarials are the basic treatment for every patient with SLE, whereas glucocorticoids should only be used when acutely indicated. If reduction

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**KEY MESSAGES**

- Systemic lupus erythematosus (SLE) is a heterogeneous disease that may affect many different organs; early diagnosis prevents adverse consequences and clearly improves the prognosis.
- The diagnosis of SLE is symptom-based and often requires an interdisciplinary approach. Depending on the organ(s) involved, patients with SLE should be referred to qualified medical specialists with long-term experience in the treatment of this disease or to a dedicated SLE center.
- The goal of treatment is remission or control of disease activity and prevention of (further) organ damage with the minimum possible dose of glucocorticoids.
- Treatment with the drugs available can clearly improve the short- and long-term prognosis of SLE. Antimalarials should be used—unless contraindicated—in all patients with SLE.
- A modern treatment strategy should comprise not only preventive measures but also the treatment of comorbidities (e.g., infections and cardiovascular events).
or tapering of glucocorticoids proves impossible, extended immunosuppression with azathioprine, methotrexate, or mycophenolate mofetil is recommended. Should the patient fail to respond, belimumab can be administered.

Although mycophenolate mofetil is not officially approved for use in SLE, there is now international consensus that its good efficacy with low toxicity make this agent an effective alternative to short-term administration of low-dose intravenous cyclophosphamide in the induction therapy for proliferative lupus nephritis.

Conflict of interest statement
Prof. Kuhn has received consultancy fees from Biogen, GlaxoSmithKline (GSK), Grüntenthal, and Lilly. She has received reimbursement of costs for conference attendance, travel, and accommodation as well as payments for the preparation of scientific meetings from Basilea, Biogen, GSK, La Roche Posay, Lilly, and Spirig Pharma GmbH.

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For eReferences please refer to:
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REVIEW ARTICLE

The Diagnosis and Treatment of Systemic Lupus Erythematosus

Annegret Kuhn, Gisela Bonsmann, Hans-Joachim Anders, Peter Herzer, Klaus Tenbrock, Matthias Schneider

eREFERENCES

**eTable**

<table>
<thead>
<tr>
<th>Subtype</th>
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| **Acute cutaneous lupus erythematosus (ACLE)** | - Localized: “butterfly rash”  
- Generalized: maculopapular exanthema  
- Oral mucous membrane: erosions, ulcers  
- Diffuse thinning of hairline (“lupus hair”) |
| **Subacute cutaneous lupus erythematosus (SCLE)** | - Annular and/or papulosquamous/psoriasiform with polycyclic confluence  
- Healing without scarring, vitiligo-like hypopigmentation  
- High photosensitivity  
- 70–90% anti-Ro/SSA and in 30–50% anti-La/SSB antibodies  
- ≥ 4 ACR criteria in 50%, development of a mild form of systemic lupus erythematosus in 10–15% (rarely involvement of kidneys and central nervous system) |
| **Chronic cutaneous lupus erythematosus (CCLE)** | - Localized (ca. 80%) or disseminated (ca. 20%)  
- Discoid erythematosus plaques with firmly adherent follicular hyperkeratoses  
- Healing with scarring (on the scalp, scarring alopecia) |
| **Discoid lupus erythematosus (DLE)** | - Tender, livid red swelling, sometimes with erosion/ulceration  
- Localization: symmetrical, cold-exposed areas of extremities |
| **Chilblain lupus erythematosus (CHLE)** | - Subcutaneous, nodular/plaque-like, dense infiltrates  
- Ulceration and calcification possible, healing with scarring and deep lipatrophy |
| **Lupus erythematosus profundus/panniculitis (LEP)** | - Erythematous, urticaria-like, edematous plaques without epidermal involvement  
- High photosensitivity  
- Variable course, healing without scarring |

*Modified after (8)