A B S T R A C T
Systemic lupus erythematosus (SLE) is a complex multi-systemic autoimmune disease with considerable clinical and immunological heterogeneity. Family physicians should be familiar with the protean manifestations of SLE to aid early diagnosis and monitoring of disease progression. The role of family physicians in SLE includes education, counselling, psychological support, management of mild disease, and recognition of the need for referral to other specialists for more serious disease and complications. Surveillance of cardiovascular risk factors and osteoporosis and advice about vaccination and reproductive issues can be performed in the primary care setting under close collaboration with rheumatologists and other specialists. This review provides family physicians with the latest classification criteria for SLE, recommendations on SLE-related health issues, and pharmacological therapies for SLE.

Introduction
Systemic lupus erythematosus (SLE) is a prototypical multi-systemic autoimmune disease that predominantly affects women of childbearing age. The disease has considerable clinical and immunological heterogeneity; no two patients with SLE are exactly alike. The pathogenesis of SLE remains obscure, with multiple genetic, epigenetic, hormonal, and immunopathological pathways being involved.1 The course of SLE is largely unpredictable and characterised by periods of disease exacerbation and remission that lead to progressive organ damage and dysfunction.2 Compared with the age- and sex-matched general population, SLE is associated with at least a five-fold increase in mortality.3 Patients with SLE have reduced quality of life because of multiple factors, such as organ damage, anxiety, and depression.4,5

In Hong Kong, the prevalence and annual incidence of SLE are estimated to be 0.1% and 6.7 per 100,000 population, respectively.6 The 15-year cumulative survival of local Chinese patients with SLE managed in non-academic hospitals is 86%.7 Infections, cardiovascular events, and malignancies are their most common causes of death. Renal and musculoskeletal complications (eg, avascular bone necrosis and osteoporotic fracture) are the most important contributors to disease and treatment-related organ damage, respectively. One-third of such patients lose their ability to work within 5 years after disease onset; this is mainly attributed to musculoskeletal pain, fatigue, anxiety/depression symptoms, and memory deterioration.8

Strategies for management of SLE by family physicians should target early recognition and diagnosis, treatment and monitoring of mild disease, and referral to specialists to formulate an individualised plan based on age, disease severity, organ function, and other medical co-morbidities.9 In this article, the latest criteria for classification of SLE, use of autoantibodies for diagnosis and assessment, the role of family physicians, disease monitoring, advice on various SLE-related health issues from a local perspective, and pharmacological treatment of the disease will be reviewed.

Classification criteria for systemic lupus erythematosus
The manifestations of SLE are protean, and any body system can be involved during the course of the disease. In our experience with 803 Hong Kong Chinese patients with SLE, the most common features are arthritis, glomerulonephritis, facial rash, and haematological disease (Fig).7 In primary care practice, the most frequently encountered early symptoms of SLE include systemic upset (fatigue, fever, weight loss, loss of appetite, and prolonged influenza-like illness), arthralgia or arthritis, facial rash, photosensitivity, mouth sores, pleuritic chest pain, and Raynaud's phenomenon.6 In hospital practice, more serious manifestations of SLE such as rapidly progressive glomerulonephritis, pulmonary haemorrhage, cardiac tamponade, severe cytopenia, and neuropsychiatric symptoms may be encountered.
The American College of Rheumatology (ACR) classification criteria for SLE were established in 1982,\(^{10}\) and this set of criteria was revised in 1997,\(^ {11}\) with the deletion of the LE cell phenomenon and the addition of antiphospholipid antibodies as a criterion. A classification of SLE is made when four or more of the 11 clinical or serological criteria are fulfilled serially or simultaneously. However, in real life practice, many patients with autoimmune cytopenia or hypocomplementaemia are treated as having SLE, even though they do not fulfil the 1997 criteria.\(^ {11}\) Moreover, dermatological manifestations other than malar rash and discoid lesions and neuropsychiatric manifestations other than psychosis and seizure are not included in the criteria. Owing to these limitations, the Systemic Lupus International Collaborating Clinics (SLICC) group revised and validated a new set of classification criteria in 2012.\(^ {12}\) A patient is classified as having SLE when at least four of the 17 SLICC/ACR criteria are fulfilled. A comparison of the 1997 ACR and 2012 SLICC criteria is shown in Table 1.

The SLICC group emphasises the absolute requirement of at least one clinical or immunologic criterion for a classification of SLE. Lupus malar rash and photosensitivity are no longer separated into two criteria. There is no need to demonstrate the absence of radiological erosion in lupus arthritis. A number of types of subacute and chronic lupus skin lesions are included, and diffuse non-scarring alopecia (excluding alopecia areata or other causes) is also regarded as a criterion for SLE classification. Haemolytic anaemia, leukopenia/lymphopenia, and thrombocytopenia are separated into three criteria, and more neuropsychiatric features are included in addition to psychosis and seizure. For the renal criteria, the dipstick test for urine protein is replaced by either the protein-to-creatinine ratio (spot urine test) or 24-hour urine protein quantification. Finally, an entity called “stand-alone” lupus nephritis is introduced, in which a patient has typical renal biopsy features of lupus nephritis and a positive ANA or anti-dsDNA antibody test in the absence of other features of SLE. The features in the SLICC criteria must be related to active SLE instead of other causes or differential diagnoses. Validation of the SLICC criteria has demonstrated higher sensitivity (97% vs 83%) but lower specificity (84% vs 96%) for SLE than the 1997 ACR criteria.\(^ {12}\)

To further improve the sensitivity and specificity of SLE classification, the ACR/EULAR (European League Against Rheumatism) is currently validating a new set of criteria consisting of an entry criterion and 10 domains (seven clinical and three immunologic). More items with different weighted scores are included. Applications on mobile devices and desktop computers will be devised to facilitate the calculation of summed scores for classification purposes.

These classification criteria for SLE are being developed to facilitate research and comparison among different cohorts of patients. Although they generally have good specificity to aid diagnosis, false positivity and negativity are bound to occur. The final diagnosis of SLE still requires the meticulous clinical judgement of attending physicians.

**Antinuclear antibody for diagnosis of systemic lupus erythematosus**

Antinuclear antibody (ANA) is the hallmark of SLE. Although this antibody shows extreme sensitivity for SLE (>98%), it has low specificity. As many as 20% to 23% of normal healthy individuals test positive...
for ANA, particularly older subjects. Other autoimmune and non-immune chronic illnesses also generate positivity for ANA, making it grossly unsuitable as a sole diagnostic test. However, ANA is an excellent screening test for SLE, and a negative result by indirect immunofluorescence assay (IIFA) may virtually exclude the diagnosis.

Antinuclear antibody is conventionally detected by the IIFA method, which involves initial screening, serial serum dilution, and determination of the distinct ANA staining patterns on human epithelial cell (HEp-2) slides. This is the most sensitive method of ANA detection, but it is labour-intensive and subject to inter-observer reading variability. Although IIFA remains the gold standard of ANA detection, automated and less laborious quantitative methods are often used by service laboratories. Enzyme-linked immunosorbent assay is commonly used to detect serum autoantibodies directed against antigens coated onto plates. As the antigens used may be derived from animal tissues or recombinant techniques, the specificity and sensitivity of the results for assessment of SLE vary among different commercial kits adopted by different laboratories. In general, higher ANA titres result in more specific predictions for SLE and related autoimmune and non-immune chronic illnesses.

**TABLE 1. Classification criteria for SLE**

<table>
<thead>
<tr>
<th>1997 ACR criteria</th>
<th>2012 SLICC/ACR criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of clinical criteria</td>
<td>11</td>
</tr>
<tr>
<td>Dermatological</td>
<td>1. Malar rash (fixed erythema, flat or raised, over the malar prominence, tending to spare the nasolabial folds)</td>
</tr>
<tr>
<td></td>
<td>2. Photosensitivity</td>
</tr>
<tr>
<td>Dermatological</td>
<td>3. Discoid lupus (typical)</td>
</tr>
<tr>
<td>Mucosal</td>
<td>4. Oral/nasal ulcers (usually painless, observed by physician)</td>
</tr>
<tr>
<td>Alopecia</td>
<td>-</td>
</tr>
<tr>
<td>Musculoskeletal</td>
<td>5. Non-erosive arthritis in ≥2 joints (tender, swollen or effusion)</td>
</tr>
<tr>
<td>Serositis</td>
<td>6. Pleurisy OR pericarditis</td>
</tr>
<tr>
<td>Haematologic</td>
<td>7. Haemolytic anaemia with reticulocytosis OR leukopenia (&lt;4000/mm³ at least twice) OR lymphopenia (&lt;1500/mm³ at least twice) OR thrombocytopenia (&lt;100/cm³)</td>
</tr>
<tr>
<td>Renal</td>
<td>8. Proteinuria &gt;500 mg/day or &gt;3+ by dipstick OR cellular cast (RBC, haemoglobin, granular, tubular, or mixed)</td>
</tr>
<tr>
<td>Neuropsychiatric</td>
<td>9. Seizure OR psychosis</td>
</tr>
<tr>
<td>No. of serological criteria</td>
<td>2</td>
</tr>
<tr>
<td>Immunologic/serological</td>
<td>1. Positive ANA (not induced by drugs)</td>
</tr>
<tr>
<td></td>
<td>2. Positive for anti-dsDNA OR anti-Sm OR antiphospholipid antibodies (IgG/IgM anticardiolipin; lupus anticoagulant; or a false positive serologic test for syphilis known to be positive for ≥6 months)</td>
</tr>
<tr>
<td></td>
<td>1. Positive ANA (above laboratory reference range)</td>
</tr>
<tr>
<td></td>
<td>2. Anti-dsDNA (above laboratory reference range or &gt;2-fold the reference range if tested by ELISA)</td>
</tr>
<tr>
<td></td>
<td>3. Anti-Sm positivity</td>
</tr>
<tr>
<td></td>
<td>4. Antiphospholipid antibody positivity: positive lupus anticoagulant, false positive result for rapid plasma regain, medium-/high-titre anticardiolipin (IgG/A/M), or anti-β₂ glycoprotein I (IgG/A/M)</td>
</tr>
<tr>
<td></td>
<td>5. Low complements (C3/C4/CH50)</td>
</tr>
<tr>
<td></td>
<td>6. Positive direct Coombs’ test (in the absence of haemolytic anaemia)</td>
</tr>
<tr>
<td>Total No. of criteria</td>
<td>11</td>
</tr>
<tr>
<td>SLE classification</td>
<td>≥4/11 Criteria</td>
</tr>
</tbody>
</table>

Abbreviations: ACR = The American College of Rheumatology; ANA = antinuclear antibody; ELISA = enzyme-linked immunosorbent assay; Ig = immunoglobulin; RBC = red blood cells; SLE = systemic lupus erythematosus; SLICC = Systemic Lupus International Collaborating Clinics
disorders. Therefore, ANA should be interpreted in the clinical context, and a diagnosis of SLE should not be based on a positive ANA result alone.

The dense fine speckle (DSF) pattern of ANA in IIFA is related to autoantibodies against a 70-kDa protein (DSF70). Interestingly, this anti-DSF70 antibody is present in around one-third of ANA-positive healthy subjects, in contrast to less than 1% of ANA-positive patients with SLE and other autoimmune diseases. Anti-DSF70 is becoming a part of the standard report along with the ANA result in public hospitals. Positive results for both ANA and anti-DSF70, in the absence of other autoantibodies such as anti-dsDNA and anti–extractable nuclear antigen (anti-ENA), may virtually exclude SLE or an ANA-related autoimmune disorder.

**Systemic lupus erythematosus diathesis recognition**

Table 2 shows a list of pointers that should alert family physicians to consider the possibility of SLE. When SLE is suspected, ANA should be included in the screening blood tests. If the patient is positive for ANA, more specific tests such as those for anti-dsDNA, anti-ENA, antiphospholipid antibodies (eg, anticardiolipin antibodies), and complements are needed to confirm the diagnosis. Other relevant investigations are also needed, such as urine analysis, cell counts, renal and liver function tests, and tests for inflammatory markers such as erythrocyte sedimentation rate or C-reactive protein (CRP). Diagnosis of SLE is based on a combination of compatible clinical features and the presence of relevant immunological abnormalities. Therefore, SLE should never be diagnosed by abnormal antibody tests alone.

The ANA titre is not useful for monitoring of SLE activity. The anti-dsDNA titre and complement levels (C3/4) are the standard serological tests for disease activity evaluation ("lupus serology"). ENAs include a number of soluble cytoplasmic and nuclear antigens. The six main antigens used to detect anti-ENA antibodies are Ro, La, Sm, RNP, Scl-70, and Jo-1. The detected antibodies are associated with certain manifestations of SLE (eg, anti-Ro with cutaneous lupus and photosensitivity) and are relevant in pregnancy (eg, anti-Ro with congenital heart blockage and neonatal lupus). Anti-Sm is specific to SLE and is a criterion for its classification. As anti-ENA antibodies seldom sero-convert over time, repeating the tests during routine follow-up is not necessary. Table 3 summarises the assessment and monitoring of patients with SLE and includes general advice about various health-related issues.

**Role of family physicians**

According to our experience with Chinese patients with SLE, mood disorders are its most frequent psychiatric manifestations. The major self-reported symptoms that lead to impaired quality of life are problems with memory and concentration and symptoms of anxiety and depression. Therefore, patients with SLE should receive education about the disease, psychological counselling, and support in the primary care setting.

In addition to understanding the clinical presentation of SLE for early diagnosis, trained family physicians are able to treat and monitor mild SLE, which comprises the following characteristics: (1) diagnosis clearly established; (2) clinically stable; (3) absence of life-threatening manifestations; (4) stable function of organ systems; and (5) absence of significant complications related to disease activity or treatment. Patients with stable SLE should be followed at intervals of 3 to 6 months. Referral to specialists is indicated for worsening disease activity, involvement of major organs such as the kidneys, haematological and central nervous system complications, development of disease or treatment-related complications, antiphospholipid syndrome, and advice about pregnancy, surgery, and other special circumstances.

Family physicians may help to monitor disease activity and the adverse effects of drug therapies. A complete blood count, renal function, SLE serology (anti-dsDNA and complements), and urinary protein analysis should be performed every 3 to 6 months for patients with stable disease. As patients with SLE are more prone to accelerated atherosclerosis as a result of disease activity and
TABLE 3. Assessment and monitoring of patients with SLE and general advice

<table>
<thead>
<tr>
<th>Initial evaluation</th>
<th>Subsequent follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>Screening by ANA in suspected cases of SLE</td>
<td>Follow-up frequency: 1-3 months, or even more frequent, for active SLE; 3-6 months for mild and stable SLE</td>
</tr>
<tr>
<td>Clinical history and physical examination for skin/scalp lesions, arthritis, lymphadenopathy, alopecia, mucosal ulceration, and Raynaud’s phenomenon. Enquire for family history of SLE or other autoimmune diseases</td>
<td>Focused history on new symptoms/manifestations. Differentiate disease activity from organ damage, drug toxicities, and co-morbidities that are causing the symptoms</td>
</tr>
<tr>
<td>Drug review, particularly those that may induce ANA/autoimmunity, eg, iononizid, methylidopa, hydralazine, minocycline, carbimazole, and phenytoin</td>
<td>Review medications and adverse effects</td>
</tr>
<tr>
<td>Vital signs, blood pressure, BMI</td>
<td>Vital signs, blood pressure, BMI, urine protein analysis</td>
</tr>
<tr>
<td>If ANA positive, check anti-dsDNA, anti-ENA, complements, anticytokelin; cell counts, liver and renal function; ESR, CRP, immunoglobulin level; urine analysis, baseline chest radiograph</td>
<td>Cell counts, renal and liver function tests, anti-dsDNA, complements, CRP (for selected patients). Regular surveillance for vascular risk factors, eg, lipid profile and glucose</td>
</tr>
</tbody>
</table>

**General advice about various health issues**

**Photoprotection**
Avoid prolonged sun exposure; protective clothing; hat and umbrella; topical sunscreen (SPF ≥30) every 1-2 hours

**Monitoring of BMD**
Baseline DXA scan; lifestyle modification plus calcium and vitamin D (if prednisolone ≥2.5 mg/day for ≥3 months); anti-osteoporotic medications if risk of major fracture ≥10% or hip fracture ≥1% at 10 years (age ≥40 years); or history of fragility fracture, BMD Z-score <-2.5 or rapid loss of ≥10% BMD in 1 year (age <40 years). Choice: oral BSP > intravenous BSP > teriparatide, denosumab, or raloxifene (not for premenopausal women); repeat DXA every 2-3 years

**Review and advice on vaccination**
Seasonal influenza and pneumococcal vaccines; hepatitis B vaccine (if at risk and not received at birth); haemophilus and meningococcal vaccines (post-splenectomy, persistent low complements); HPV vaccine (age >9 years); avoid live attenuated vaccine but consider herpes zoster vaccine when not receiving intensive immunosuppression; vaccination should be given during period of disease remission and minimal immunosuppression

**Counselling on pregnancy and assisted reproduction**
Conception may be considered when disease is in remission for ≥6 months; adjustment of medications to avoid teratogenicity; check anti-Ro and aPL antibodies; ovulation stimulation may be done when disease is in remission for ≥6 months; aspirin and heparin prophylaxis if aPL antibodies positive but no history of thrombosis or other contra-indications

**Counselling on contraception**
Barrier method; progestogens; progestogen-impregnated IUD; low-dose oestrogen containing combined OC pill is not contra-indicated for stable and quiescent disease and in the absence of aPL antibodies or other contra-indications

**Abbreviations:** ANA = antinuclear antibody; aPL = antiphospholipid; BMD = bone mineral density; BMI = body mass index; BSP = bisphosphonate; CRP = C-reactive protein; DXA = dual-energy X-ray absorptiometry; ENA = extractable nuclear antigen; ESR = erythrocyte sedimentation rate; HPV = human papillomavirus; IUD = intrauterine device; OC = oral contraceptive; SLE = systemic lupus erythematosus; SPF = sun protection factor

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**Advice on photoprotection**

The ultraviolet (UV) light spectrum can be divided into UVC (100-290 nm), UVB (290-320 nm), and UVA (320-400 nm) wavelengths. The superficial layers of the epidermis mainly absorb UVB irradiation, but longer-wavelength UVA can also penetrate the deeper dermis. Ultraviolet light may trigger a complicated process that includes the activation of keratinocytes to release pro-inflammatory cytokines, chemokines, and interferons, which may exacerbate local and systemic autoimmunity.25

Photosensitivity was poorly described in the ACR criteria as skin rash resulting from an unusual reaction to sunlight, as reported in patients’ history or physicians’ observation.10,11 Some clinicians regard photosensitivity as induction or exacerbation of skin lesions after extensive sun exposure, which also includes sunburn. Because of the broad definition of photosensitivity, its incidence in SLE ranges widely (27%-100% in different studies).25 The latency period between UV exposure and skin eruptions can range from several days to 3 weeks. In addition to UV exposure, photosensitivity in SLE can be caused by photosensitising medications and co-existing photodermatosis.

Avoidance of excessive sunshine, particularly during midday hours, is often advised to patients with SLE. Hats, protective clothing, and umbrellas are effective at blocking UV light. Ultraviolet-
protective sunglasses and lip balms may also help. Topical sunscreen is a common means of reducing UV light penetration. A sunscreen with sun protection factor 30 absorbs/reflects 97% of UV light.30 Patients with SLE should apply sunscreen (sun protection factor ≥30) 30 minutes before going out into the sun to all exposed body parts and re-apply it after 1 to 2 hours if exposure is to continue. Patients should be reminded that sunscreen does not provide 100% protection from UV light or offer skin support for repair of photodamage. Therefore, avoidance of unnecessary sun exposure remains the most important behavioural modification.

Vitamin D supplementation and osteoporosis prevention

Vitamin D deficiency has recently been postulated to be an environmental trigger for autoimmune diseases, including SLE.27 Compared with age- and sex-matched healthy subjects, patients with SLE have significantly lower serum vitamin D levels, which correlate inversely with disease activity.28-30 Vitamin D insufficiency in SLE has multiple contributing factors, which include avoidance of UV exposure by using sunscreen, chronic kidney disease, long-term use of medications that hamper absorption or metabolism of vitamin D, and anti-vitamin D antibodies that may enhance plasma clearance of vitamin D.27 Although there is conflicting evidence regarding the efficacy of vitamin D supplementation at alleviating clinical SLE activity, such supplementation is recommended for prevention and treatment of glucocorticoid-induced osteoporosis.31 According to the updated ACR recommendations, patients receiving ≥3 months of prednisolone (≥2.5 mg/day) should receive elemental calcium (1000-1200 mg/day) and cholecalciferol (600-800 IU/day) along with lifestyle modification (weight bearing exercise, cessation of smoking, balanced diet, and maintaining optimal body weight).31 In patients with SLE aged >40 years, who have a moderate to high risk of a major osteoporotic (>10%) or hip fracture (>1%) within 10 years (as assessed by the fracture risk assessment tool), oral bisphosphonates are recommended. When oral bisphosphonates are inappropriate (eg, owing to intolerance or contra-indication), intravenous bisphosphonates (eg, zoledronate) are the next alternatives to be considered. Other treatment options include teriparatide (which is costly and inconvenient to inject daily), denosumab, and raloxifene (which has a lack of efficacy data regarding fractures). There is a general paucity of efficacy data of these agents in younger patients aged <40 years. The ACR recommends treatment for moderate–to-high-risk younger patients, defined as having a previous osteoporotic fracture; bone mineral density Z-score of <−3.0 at the hip or spine; or rapid loss of ≥10% bone mineral density over 1 year and continuous prednisolone treatment (≥7.5 mg/day for ≥6 months).31 The choice of drugs is the same as that for older patients, except for raloxifene, which is not indicated in premenopausal women or male patients.

Vaccination

Patients with SLE are prone to infections because of the underlying immune aberrations and therapies with immunosuppressive regimens.1 Vaccination offers the most cost-effective method of reducing infection risk in patients with SLE. Non-live vaccines such as influenza and pneumococcal vaccines are generally well tolerated in SLE, although they are less immunogenic than in age-matched individuals.32 Although there is conflicting evidence on whether influenza vaccine exacerbates SLE activity,33 seasonal influenza vaccination according to national guidelines is recommended.34,35 Influenza and pneumococcal vaccination is particularly recommended for patients with SLE before rituximab therapy. Additional vaccinations against *Haemophilus influenzae* and *Neisseria meningitidis* are suggested for patients with functional asplenia, splenectomy, or persistently very low complement levels.34,35 Hepatitis B vaccination can be safely administered to patients with SLE who are at risk of infection if it was not given at birth. Female patients with SLE are more prone to persistent genital human papillomavirus (HPV) infection, which predisposes them to cervical cancers. The HPV vaccine is recommended for patients with SLE, preferably prior to the beginning of sexual activity. There is no evidence of increased SLE flares after administration of the quadrivalent or bivalent HPV vaccines.36 In Hong Kong, the quadrivalent and nonavalent HPV vaccine is licensed for patients aged >50 years. Patients with SLE who are at risk of cervical cancers. The HPV vaccine is recommended for patients with SLE during periods of disease quiescence and minimal immunosuppression.

Live attenuated vaccines are generally not recommended for individuals who are heavily immunocompromised because of the risk of disseminated infections. Of relevance is the live attenuated herpes zoster vaccine, which has been licensed for patients aged ≥50 years. Patients with SLE are particularly prone to herpes zoster reactivation, with a pooled relative risk of 2.10 compared with the age- and sex-matched general population.38 According to the United States Advisory Committee on Immunization Practices, herpes zoster vaccine should not be given to individuals who are receiving heavy immunosuppressive therapies, such as prednisolone (>20 mg/day for ≥2 weeks), methotrexate (≥0.4 mg/kg/week), and azathioprine (≥3.0 mg/kg/day).39 However, in view of the high incidence of herpes zoster in patients with SLE, herpes zoster vaccine should be considered in those
who have stable and remitted disease that does not require intense immunosuppression.\textsuperscript{34,35} The herpes zoster vaccine has been administered safely to SLE patients without subsequent development of herpetiform lesions or disease flares.\textsuperscript{40}

**Pregnancy counselling, assisted reproduction, and contraception**

The fertility of patients with SLE is preserved, unless they develop chronic kidney disease or have been treated with cyclophosphamide. Patients with SLE should not be discouraged regarding pregnancy, provided that their disease has been under good control for at least 6 to 12 months.\textsuperscript{41} The outcomes of pregnancies have improved for patients with SLE in the past few decades as a result of better risk stratification, pre-conception counselling, and close multidisciplinary surveillance. However, the rates of pregnancy loss, preterm birth, pre-eclampsia, and intrauterine growth retardation remain higher in pregnancies of patients with SLE than in those of patients without.\textsuperscript{41} The main risk factors for poor maternal and fetal outcomes in pregnancies of patients with SLE are active disease at conception (particularly nephritis), the presence of strongly positive antiphospholipid antibodies (or a history of obstetric antiphospholipid syndrome), and a history of lupus nephritis.\textsuperscript{42} Some medications such as cyclophosphamide, mycophenolate mofetil, leflunomide, and angiotensin-converting enzyme inhibitors/angiotensin receptor blockers are teratogenic. High-dose glucocorticoid treatment may lead to intrauterine growth retardation and premature delivery. The risk of congenital heart blockage in anti-Ro-positive mothers with SLE is approximately 1% to 2%.\textsuperscript{42} Close liaison with obstetricians and paediatricians for monitoring of the cardiovascular status of the fetus during pregnancy and assessment of neonatal lupus syndrome is needed. In general, SLE patients with ≥6 months of disease remission who are in good general health may consider conception. Referral to specialists for adjustment of medications and prophylactic heparin/asaarin (in case of obstetric antiphospholipid syndrome) is needed.

The use of assisted reproductive technology is increasing. Despite increases in disease flares and thrombosis after hormonal ovulation stimulation,\textsuperscript{43} the current recommendation is to individualise the risk of these procedures in patients with SLE.\textsuperscript{44} Assisted reproductive technology procedures should be discouraged in female SLE patients who have active disease, severe renal insufficiency, serious valvulopathy or coronary heart disease, poorly controlled hypertension, history of major thrombotic events, or antiphospholipid syndrome.\textsuperscript{44} Counselling should also be given about other serious adverse effects of assisted reproductive technology procedures, such as ovarian hyperstimulation. In patients with SLE who are positive for antiphospholipid antibodies and have no history of thrombosis, aspirin and heparin prophylaxis is recommended during these procedures.\textsuperscript{45} Similar to naturally achieved pregnancies, the SLE of candidates for assisted reproductive technology should have been quiescent for ≥6 months.\textsuperscript{46}

Patients with SLE should be counselled about contraception methods. Barrier methods are generally safe. Oestrogen-containing oral contraceptive pills were discouraged in the past. However, in a randomised double-blind placebo-controlled trial, a combination of oral contraceptive pills was not shown to increase SLE disease flares or thrombosis after 12 months’ administration as compared with placebo in patients with stable SLE and no antiphospholipid antibodies.\textsuperscript{47} Another randomised controlled trial did not reveal a difference in disease flares or adverse events in 12 months among patients with SLE who were assigned to receive combined oral contraceptive pill, intrauterine device, and progestogen-only pills for contraception.\textsuperscript{48} Thus, patients with stable SLE and no antiphospholipid antibodies or other contra-indications may use low-dose oestrogen oral contraceptive pills if they want to adopt a more reliable contraceptive method. When oral contraceptive pills are not appropriate, progestogens and intrauterine devices can be offered to patients with SLE as alternatives.\textsuperscript{44} Progestogen-impregnated intrauterine devices have the advantage of reducing the incidence of dysmenorrhoea and irregular vaginal bleeding.\textsuperscript{44}

**Conventional and novel therapeutics for systemic lupus erythematosus**

Hydroxychloroquine is an antimalarial drug that exhibits immune-modulatory properties in addition to antithrombotic and lipid- and glucose-lowering properties.\textsuperscript{49} Hydroxychloroquine is mainly indicated for skin, joint, and serosal manifestations of SLE and has a glucocorticoid-sparing effect. The drug is compatible with pregnancy and breastfeeding and is relatively safe to be prescribed and monitored by trained family physicians. Allergy and acute ocular and neuromuscular toxicity are rare adverse drug reactions. Chronic use of hydroxychloroquine may lead to retinopathy, with the main risk factors being older age, pre-existing liver and renal dysfunction, higher daily dose, and longer duration of therapy.\textsuperscript{50,51} Early recognition of this adverse drug reaction is essential to minimise damage to vision. Referral to an ophthalmologist for baseline examination and regular retinopathy surveillance is recommended.\textsuperscript{50}

In a recent study, 2361 patients received
hydroxychloroquine for >5 years. In that study, the risk of retinopathy was <1% in the first 5 years and <2% in 10 years when the daily dose was <5 mg/kg of real body weight. The risk of retinopathy increased sharply to 20% after 20 years. The daily dose of hydroxychloroquine was the most critical factor for the retinopathy risk, which correlated better with real rather than ideal body weight. The American Academy of Ophthalmology recommends a maximum daily hydroxychloroquine dose of <5.5 mg/kg of real weight to minimise retinal toxicity. A baseline ophthalmologic examination within the first year of commencement of drug administration is recommended, and annual screening should start after 5 years of exposure in patients using a lower dosage and without major risk factors. Patients with major risk factors for retinopathy (older age, renal or liver dysfunction, or pre-existing macular or retinal disease) should be screened annually if not more frequently.

Short courses of non-steroidal anti-inflammatory drugs (NSAIDs) are indicated for control of SLE symptoms such as arthritis, myalgia, serositis, and fever. The risk of allergic and skin reactions, aseptic meningitis, and renal and liver toxicity is increased in SLE patients, despite their younger age. Ovulation may be affected by NSAIDs, and they should be used cautiously during pregnancy. Patients with SLE who have renal insufficiency, bleeding tendency, and pre-existing coronary heart disease should avoid NSAIDs. Except for their lower risk of gastrointestinal toxicity, selective Cox II inhibitors share similar renal, hepatological, and neurological adverse effects with non-selective Cox inhibitors. Among the NSAIDs, naproxen appears to be associated with the lowest risk of cardiovascular toxicity.

<table>
<thead>
<tr>
<th>Medication</th>
<th>Indication</th>
<th>Dosage</th>
<th>Monitoring and precaution</th>
</tr>
</thead>
<tbody>
<tr>
<td>HCQ</td>
<td>Skin and joint disease, lupus nephritis, and general use to reduce organ damage</td>
<td>200-400 mg/day, keep ≤200 mg/day for maintenance to reduce retinopathy risk</td>
<td>Baseline ophthalmological examination and regular surveillance</td>
</tr>
<tr>
<td>NSAIDs</td>
<td>Arthritis, fever, serositis</td>
<td>Depends on preparation</td>
<td>Avoid in chronic kidney disease, history of arterial thrombosis, and multiple vascular risk factors</td>
</tr>
<tr>
<td>Glucocorticoids</td>
<td>Low dose for mild disease and maintenance; moderate to high dose for major organ disease, eg, nephritis, nephropathy or myelopathy</td>
<td>Low dose: &lt;10 mg/day Moderate dose: 10 mg/day - 0.5 mg/kg/day High dose: ≥0.5-1.0 mg/kg/day</td>
<td>Excludes infection before administration; monitoring of blood pressure, lipid and glucose level; prevention of osteoporosis and infections</td>
</tr>
<tr>
<td>Methotrexate</td>
<td>Arthritis not responding to HCQ, myositis, skin lupus, serositis</td>
<td>7.5-15 mg/day</td>
<td>May aggravate photosensitivity; monitoring of liver/renal function, cell counts, chest radiograph and lung function; contra-indicated in severe renal insufficiency</td>
</tr>
<tr>
<td>Azathioprine</td>
<td>Lupus nephritis, more serious organ disease, glucocorticoid sparing effects</td>
<td>1-2 mg/kg/day</td>
<td>Close monitoring of leukocyte and neutrophil counts when first used; regular blood counts and liver function</td>
</tr>
<tr>
<td>MMF</td>
<td>Lupus nephritis, refractory skin and more serious organ disease</td>
<td>1-3 g/day</td>
<td>Monitoring for gastrointestinal intolerance and cell counts</td>
</tr>
<tr>
<td>CSA/TAC</td>
<td>Lupus nephritis, refractory haematological and more serious organ disease</td>
<td>TAC: 0.06-0.1 mg/kg/day CSA: 3-5 mg/kg/day</td>
<td>Monitoring of blood pressure, electrolytes, renal function, and symptoms of neurotoxicity</td>
</tr>
<tr>
<td>CYC</td>
<td>Lupus nephritis, neuropsychiatric, and other serious major organ diseases</td>
<td>Oral: 1-2 mg/kg/day Intravenous pulse: 0.5-1 g/m² per pulse</td>
<td>Oral regimen no more than 3 months; pulse regimen no more than 6 pulses, unless neuropsychiatric or more serious manifestations; monitoring of leukocyte and neutrophil counts and cytostasis; prophylaxis against PCP in selected patients</td>
</tr>
<tr>
<td>Belimumab</td>
<td>Moderate to severe SLE manifestations not responding to standard of care</td>
<td>10 mg/kg intravenous at baseline, week 2, week 4, and then 4-weekly</td>
<td>Not indicated for serious nephritis, neuropsychiatric and other serious organ manifestations; monitor cell counts and for infusion reaction</td>
</tr>
<tr>
<td>Rituximab</td>
<td>Refractory SLE manifestations, particularly lupus nephritis</td>
<td>Intravenous: 375 mg/m² weekly for 4 doses or 1 g 2-weekly for 2 doses (one course)</td>
<td>Prophylaxis for occult hepatitis B reactivation; monitoring of immunoglobulin level and cell counts</td>
</tr>
</tbody>
</table>

**Abbreviations:** CSA = cyclosporine A; CYC = cyclophosphamide; HCQ = hydroxychloroquine; MMF = mycophenolate mofetil; NSAIDs = non-steroidal anti-inflammatory drugs; PCP = pneumocystis pneumonia; SLE = systemic lupus erythematosus; TAC = tacrolimus
events and is the preferred NSAID for patients with multiple cardiovascular risk factors. Diclofenac is associated with the highest risk and should be avoided in these patients. A recent randomised controlled trial reported that celecoxib was non-inferior to ibuprofen or naproxen with regard to cardiovascular safety in patients with rheumatoid arthritis and osteoarthritis. The lowest effective dose of NSAIDs should be used, and their indications should be periodically reviewed. Monitoring of fluid status, kidney function, liver transaminases, and blood pressure is necessary.

Glucocorticoids and a number of non-glucocorticoid immunosuppressive agents are often used to treat more serious organ manifestations of SLE. Systemic glucocorticoids are a major cause of treatment-related organ damage in patients with SLE and contribute significantly to mortality and co-morbidities. Therefore, the use of systemic glucocorticoids in SLE has to be fully justified, judicious, and closely monitored. Other treatment modalities used for severe SLE include intravenous immunoglobulin and plasmapheresis. A biological agent called belimumab has recently been approved for mild to moderate SLE manifestations that are refractory to standard therapies. Although rituximab has not been proven to be more effective than placebo in randomised controlled trials, it is often used off-label for refractory lupus manifestations. Many other biological and targeted synthetic agents are being tested in patients with SLE. While it is outside the scope of this review to describe these therapies in detail, they are summarised in Table 4 for quick reference.

Conclusions
Systemic lupus erythematosus is a prototypical autoimmune disease that affects primarily young women of reproductive age. The new SLICC classification has expanded the clinical and serological criteria for its classification. Systemic lupus erythematosus should never be diagnosed based solely on positive test results for antibodies, particularly ANA, which is highly non-specific and should be interpreted in conjunction with clinical signs and symptoms. In view of the disease’s multi-systemic involvement, holistic care is necessary to formulate treatment plans for individual patients. Family physicians play an important role in establishing an early diagnosis, treatment and monitoring of mild disease, and making referrals to specialists when appropriate. Education, counselling, and psychological support are equally important to improve treatment adherence and alleviate mood symptoms. General advice about photoprotection, vaccination, prevention of osteoporosis, and reproductive issues may be given in the primary care setting. Hydroxychloroquine is a relatively safe drug that can be commenced and monitored by family physicians. For patients with stable SLE, screening for cardiovascular risk factors and osteoporosis may also be performed periodically in family clinics.

Author contributions
The author has made substantial contributions to the concept or design, acquisition of data, analysis or interpretation of data, drafting of the article, and critical revision for important intellectual content.

Declaration
The author has disclosed no conflicts of interest. The author had full access to the data, contributed to the study, approved the final version for publication, and take responsibility for its accuracy and integrity.

References


