The Many Faces of Lupus: An Approach to the Assessment of a Lupus Patient

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Abstract  One of the first steps in evaluating a patient with lupus is to recognize that there are various subtypes of lupus. Determining which variant of lupus in afflicting the patient will determine the extent of disease involvement and even prognosis for the patient. Certain anti-nuclear antibodies and biomarkers for lupus have been shown to correlate with disease activity. Recognizing the prognostic indicators of active lupus can help guide a clinician in choosing the appropriate pharmacologic treatments for their lupus patients. This clinical review and summary of the literature was derived from medical and online databases such as PubMed, Cochrane review, up-to-date, the American College of Rheumatology and American College of Physicians Visual Diagnostic Logical Imaging resources. The following is an evidence-based, objective and balanced summation of this literature review along with the author's own rheumatologic clinical experience over the past 25 years. For the first time in decades, consensus guidelines have been established for the management of lupus nephritis. In addition, the first Food and Drug Administration-approved biologic agent, belimumab, is now available for the treatment of certain manifestations of systemic lupus erythematosus. The reader will be able to recognize at the bedside the various subtypes of lupus, parameters of disease activity and the appropriate selection of treatment modalities.

Keywords  Antiphospholipid, Autoantibodies, Lupus

1. Introduction

The word lupus means “wolf” in Latin. Historically, physicians often thought the facial lesions of lupus patients resembled the bites of a wolf. The images of this disease often raised fear and frustration among patients as well as their doctors. Over the past century, the recognition and classification of various subtypes of lupus, the fractionation of auto-antibodies and their prognostic significance have facilitated the recognition and management of the various subsets of lupus. For the first time in several decades, a biologic agent approved by the U.S. Food and Drug Administration has been added to the treatment armamentarium for systemic lupus erythematosus (1).

2. First Steps in the Evaluation of a Lupus Patient

One of the first steps in the evaluation of a patient is to recognize which population of patients is likely to succumb to a disease. Lupus is a prototypical autoimmune disease which has a female preponderance. Peak incidences occur during one’s most productive years of life: 15 through 45 years of age (2). As with many diseases, Hispanics and African-Americans are afflicted more often than Caucasians and with more severe disease (3). The pathogenesis of this disease has been recognized as probably being multi-factorial, with a genetic predisposition triggered by hormonal and environmental factors. The Major Histocompatibility Complex (MHC) Human Leucocyte Antigen (HLA) DR 2, DR 3 and B8 alleles serve as antigenic determinants, known as epitopes, contributing to the formation of pathogenic auto-antibodies due to dysregulation of self-tolerance, apoptosis and cellular inflammation (4).

The next step is to be alert for the environmental triggers that generally pre-date and/or accompany the disease presentation by performing a thorough review of systems. Antecedent infections, stress, hormonal therapies, sunlight and certain medications have been recognized as inciting triggers of lupus (5). You will learn in this monograph how certain triggers will lead to a specific type of lupus presentation, e.g. exposure to procainamide can lead to a drug-induced lupus (6).

3. Subtypes or Variants of Lupus

There are numerous subtypes of lupus as shown in Table 1 (7).
CD4 helper cell activity with sub-optimal CD8 suppressor genetic predisposition does exist with a 24-30% highly complex. Predominantly, there is an up-regulation of to SLE other than genetics (9). The pathogenesis of SLE is variation in hormones at that time (9). As mentioned earlier, incidence of 15 and 45 yrs of age is possibly due to the African-Americans and Hispanics (8). The bimodal peak system as well as the skin and joints. The female: male ratio such as the kidney, lungs, heart, brain and hematopoietic systemic form, which generally involves multiple organs leading to specific clinical manifestations. For example, anti-ds DNA antibodies have a propensity to bind to the glomerulus with activation of complement causing sub-epithelial and/or sub-endothelial immune complex deposition & significant renal damage (12). Lupus nephritis can be difficult to manage, but for the first time in several decades, members of a task force panel from the American College of Rheumatology have issued guidelines in the management of these patients (Table 3) (13).

### 3.1. Systemic Lupus Erythematosus (SLE)

The most severe and widespread form of lupus is the systemic form, which generally involves multiple organs such as the kidney, lungs, heart, brain and hematopoietic system as well as the skin and joints. The female: male ratio is 8:1, with a predisposition in minority groups such as African-Americans and Hispanics (8). The bi-modal peak incidence of 15 and 45 yrs of age is possibly due to the variation in hormones at that time (9). As mentioned earlier, genetic predisposition does exist with a 24-30% concordance among monozygotic twins, 2-5% concordance among dizygotic twins suggesting other factors contributing to SLE other than genetics (9). The pathogenesis of SLE is highly complex. Predominantly, there is an up-regulation of CD4 helper cell activity with sub-optimal CD8 suppressor cell function, over-expression of B cells with proliferation of auto-antibodies, which in turn, form immune complexes depositing on tissues causing cellular injury (10). SLE has a myriad of presentations ranging from mild, localized disease to severe multi-organ involvement abruptly or sequentially over the course of months to even years. This poses a challenge to practitioners as SLE can be a great mimic of many diseases. The 1997 American College of Rheumatology revised criteria for the classification of Systemic Lupus Erythematosus offers a list of the most typical clinical features found in this systemic form (Table 2) (11).

### Table 1. Subtypes of Lupus & Lupus-Related Syndromes*

<table>
<thead>
<tr>
<th>1.</th>
<th>Systemic Lupus Erythematosus (SLE)</th>
</tr>
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<tbody>
<tr>
<td>2.</td>
<td>Lupus Anti-coagulant (LAC) &amp; Anti-Phospholipid Antibody (APL)</td>
</tr>
<tr>
<td>3.</td>
<td>Neonatal Lupus</td>
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<td>4.</td>
<td>Drug-induced Erythematosus</td>
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<tr>
<td>5.</td>
<td>Discoid Lupus Erythematosus (DLE)</td>
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<tr>
<td>6.</td>
<td>Subacute Cutaneous Lupus Erythematosus (SCLE)</td>
</tr>
</tbody>
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Although the usual constitutional symptoms such as fatigue, malaise, some weight loss and changes in appetite can be seen with all of these variants, each subtype can present with their own specific clinical manifestations and prognostic significance.

### Table 2. 1997 American College of Rheumatology Revised Criteria for the Classification of Systemic Lupus Erythematosus*

<table>
<thead>
<tr>
<th>M</th>
<th>1. Malar Rash</th>
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<tbody>
<tr>
<td>D</td>
<td>2. Discoid Rash</td>
</tr>
<tr>
<td>S</td>
<td>3. Serositis: pleuritis, pericarditis</td>
</tr>
<tr>
<td>O</td>
<td>4. Oral Ulcers: usually painless</td>
</tr>
<tr>
<td>A</td>
<td>5. Arthritis: non-erosive</td>
</tr>
<tr>
<td>P</td>
<td>6. Photosensitivity</td>
</tr>
<tr>
<td>B</td>
<td>7. Blood disorders: low WBC, Lymphs, Hemolytic anemia</td>
</tr>
<tr>
<td>R</td>
<td>8. Renal Disease: proteinuria, RCC casts</td>
</tr>
<tr>
<td>A</td>
<td>9. Anti-ANNA: high titters of Anti-Nuclear Antibody</td>
</tr>
<tr>
<td>I</td>
<td>10. Immunologic: + Anti-ds DNA, smith, Anti-cardiolipin (ACLI) abs</td>
</tr>
<tr>
<td>N</td>
<td>11. Neurologic: seizure or psychosis</td>
</tr>
</tbody>
</table>


Although the criteria are often used for academic research and not as a sole diagnostic tool, it is still a helpful resource to implement at the bedside. Students in the medical profession often use the mnemonic “MD SOAP BRAIN” to facilitate the memorization of each item.

Often, the clinical presentation is a young female who presents with a viral or flu-type syndrome triggered by environmental factors with a photosensitive skin rash. This will often progress to internal organ involvement if there is a delay in diagnosis.

One of the most common organs involved in SLE is the kidney. It has been recognized that certain auto-antibodies have a predilection for various organs leading to specific clinical manifestations. For example, anti-ds DNA antibodies have a propensity to bind to the glomerulus with activation of complement causing sub-epithelial and/or sub-endothelial immune complex deposition & significant renal damage (12). Lupus nephritis can be difficult to manage, but for the first time in several decades, members of a task force panel from the American College of Rheumatology have issued guidelines in the management of these patients (Table 3) (13).

### Table 3. American College of Rheumatology First-Time Guidelines for Lupus Nephritis: Biopsy All*

| Active disease based on urine sediment, proteinuria, Creatinine, Complement & Anti-ds DNA antibody levels. |
| Maintain BP <130/80. |
| Statin therapy for patients with LDL cholesterol levels above 100 mg/dL. |
| Pregnancy counseling for fertile women. |
| Modification of Induction therapy for Class III/IV nephritis. |
| Refractory Patients may need biologic agent (not FDA approved but trials are currently in place). |

HCl= Hydroxychloroquine
ACEI= Angiotensin Converting Enzyme Inhibitor


In short, the take home message is to be aggressive and biopsy all lupus patients that present with nephritis, even if there is no significant change in creatinine clearance or urine sediment. Therefore, any urinalysis that demonstrates even a mild presence of microscopic hematuria or trace proteinuria without any significant decline in glomerular filtration or creatinine clearance should prompt a clinician to pursue a renal biopsy. In essence, these patients with early urine changes along with hypocomplementemia and elevated anti-dsDNA antibodies have nephritis until proven otherwise. The earlier a patient with lupus nephritis is identified, the more reversible the condition and the better the prognosis. Once a renal biopsy is obtained, there are 6 distinct histologic categories of lupus nephritis as set forth by several organizations such as the World Health Organization (WHO), International Society of Nephrology (ISN) and the Renal Pathology Society (RPS) (Table 4) (14).
Table 4. WHO & ISN/RPS Classification of Lupus Nephritis*  

<table>
<thead>
<tr>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>I. Minimal change- mild, good prognosis</td>
<td></td>
</tr>
<tr>
<td>II. Mesangial- interstitium- variable prognosis</td>
<td></td>
</tr>
<tr>
<td>III. Focal Proliferative- spotty glomeruli</td>
<td></td>
</tr>
<tr>
<td>IV. Diffuse Proliferative – aggressive</td>
<td></td>
</tr>
<tr>
<td>V. Membranous – (basement membrane) heavy proteinuria (nephritic)</td>
<td></td>
</tr>
<tr>
<td>VI. Advanced Sclerosis- end stage</td>
<td></td>
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</tbody>
</table>

WHO= World Health Organization  
ISN= International Society of Nephrology  
RPS= Renal Pathology Society  


The consensus from these organizations is that a large number of affected glomeruli with diffuse architectural destruction will most likely lead to a negative response to therapy (15, 16).

Other organ systems affected are the hematologic system, in which patients present with an array of blood disorders such as anemias, thrombocytopenias, pancytopenias, hemolytic anemias and clotting disorders; gastrointestinal involvement such as dyspepsia, autoimmune hepatitis, pancreatitis and vasculitis or ischemia of the intestine; pulmonary manifestations such as pleurisy, serositis, pneumonitis and pulmonary infarction (17).

Although not typically affected in lupus patients, the myocardium may also be involved. Libman-Sacks endocarditis (named after the scientists who discovered this sterile form of valvular abnormality) is the most characteristic cardiac manifestation in systemic lupus erythematosus. Lesions are usually clinically silent, but when such disease does occur, cardiac failure, endocarditis and embolic phenomena can cause severe neurologic and systemic complications (18).

Neuropsychiatric manifestations of SLE, although not frequent, can present with a wide range of signs and symptoms. From headache and mood disorders to active, focal neurologic deficits such as transverse myelitis, stroke and cognitive decline have all been reported (19). However, due to the difficulty in the diagnosis of these conditions independent from a lupus disorder, only seizures and psychosis are part of the SLE criteria mentioned earlier.

Fortunately, there are biomarkers that correlate with disease activity, help differentiate which subtype of lupus is involved and also serve as prognostic indicators. The anti-neuronal antibodies and anti-ribosomal antibodies correlate with CNS involvement, anti-ribosomal antibodies predisposing to Mixed Connective Tissue Disease (MCTD) and Raynaud’s phenomenon, anti-SSA(Ro) and anti-SSB(La) antibodies associated with a sicca complex as in Sjogren’s syndrome and anti-phospholipid antibody or lupus anticoagulant associated with thrombophilia (20,21).

3.2. Lupus Anti-coagulant and the Anti-Phospholipid Antibody Syndrome (LAC or APL)

Contrary to its name, the lupus anticoagulant (LAC) is actually an immunoglobulin with thrombotic properties binding to the phospholipids of cell membranes as well as plasma cell proteins such as beta 2 glycoprotein I. It is an inhibitor globulin which causes an increase in activated partial thromboplastin time when testing a patient’s serum with LAC. This and other anti-phospholipid (APL) antibodies cause a hypercoagulable state resulting in frequent miscarriages, arterial and venous thrombosis and thrombocytopenia (22). APL antibody can exist by itself and often these patients will have a history of recurrent spontaneous abortions as the sole complaint (23). A typical rash, livedo reticularis, is characteristic in these patients (Figure 1).

![Livedo Reticularis](Figure 1)

Up to 30 to 35% of SLE patients have these APL antibodies and, in some cases, a catastrophic form causing severe thrombotic strokes and even death have been reported (24). Most patients will be controlled with the use of a baby aspirin and/or warfarin. Aggressive anticoagulation and pulse steroids may be required for more severe cases.

3.3. Neonatal Lupus

Neonatal lupus results from the vertical transmission of the mother’s auto-antibodies to the fetus. The SSA/Ro, sometimes SSB/La, IGG antibodies travel across the placenta and disseminate throughout fetal tissue. Serious manifestations can occur when these auto-antibodies bind to fetal cardiac tissue and cause congenital heart block with third degree heart block, bradycardia and/or myocarditis. Thrombocytopenia and leukopenia may also occur. The rash is the least worrisome clinical manifestation, which often resolves as the maternal antibodies wane from the infant in approximately 6 to 8 months (Figure 2). Treatment applies to symptoms present (25).
3.4. Drug-induced Lupus Erythematosus

Many patients have reactions to various medications and can even present with auto-immune complications. The pathogenesis of drug-induced lupus erythematosus is not well understood. However, a common theory is that certain drugs act as hapten due to their oxidative metabolites, trigger the formation of auto-antibodies against histone proteins and cause clinical manifestations from mild, localized disease (serositis, pleuritis) to a diffuse, systemic form of lupus (26). Although there are numerous medications causing the formation of anti-histone antibodies and thus, drug-induced lupus, definitive drug causes have been identified as procainamide, minocin, hydralazine and biologics such as etanercept (27). Treatment is focused on removal of the drug and supportive care until symptoms resolve.

Steroids are rarely necessary.

3.5. Discoid Lupus Erythematosus (DLE)

Mucocutaneous forms of lupus cause significant cosmetic difficulties for patients. Most of these cutaneous forms are localized to the face, cheeks, upper arms and chest. There is tremendous variability of the rashes, ranging from the classic butterfly rash seen in SLE to fixed lesions which can scar and disfigure if not treated promptly as in Discoid Lupus Erythematosus (Figure 3).

DLE is mostly localized to the skin and rarely generalizes or evolves to a systemic form of lupus. Diagnosis is confirmed by a skin biopsy revealing hyperkeratosis, follicular plugging and immune mononuclear cells at the dermal-epidermal junction (28). The Anti-nuclear antibody (ANA) so characteristically sensitive in the detection of lupus tends to be negative in this variant. Sunscreen protection, intra-lesional steroid injections and/or anti-malarials are the usual modalities of treatment for DLE.

3.6. Subacute Cutaneous Lupus Erythematosus (SCLE)

Subacute Cutaneous Lupus Erythematosus (SCLE) is another form of mucocutaneous lupus. This cutaneous form of lupus is more common in Caucasian females. The papulosquamous rash tends to be annular, circular and non-scarring (Figure 4).

It is easily aggravated by sunlight and most often occurs on sun-exposed surfaces of the body such as shoulders, extensor surfaces of the arms, upper chest, back and neck. Clinical manifestations tend to be serositis and pleuritis with the absence of internal organ problems such as renal or cardiovascular complications. Typically, antibodies to the ribonucleoproteins SSA/Ro and SSB/La are present (29). Treatment is usually photoprotection and anti-malarials such as hydroxychloroquine.

4. Monitoring Parameters for the Management of Lupus
As in most chronic diseases, a thorough history and physical examination are paramount for the management of lupus. Educating the patients about the early signs of a lupus flare can also allow for acute intervention and prevention of more serious disease. Flare-ups of lupus can present as worsening fatigue, low-grade fevers, arthralgias, myalgias and other constitutional symptoms. The butterfly rash may appear more intense and/or other rashes may become more disseminated. An antecedent infection, recent drug use or a photosensitivity reaction can often trigger these flares (5). This should prompt the practitioner to obtain a panel of acute phase reactants and biomarkers. The acute phase reactants such as sedimentation rate and C-reactive protein usually correlate with disease activity. There have been some studies suggesting the high sensitivity C-reactive protein (hs-CRP) has been used as a risk assessment for cardiovascular disease in lupus patients (30). Other serologic biomarkers such as elevated anti-ds DNA antibodies and low complement levels such as CH50, C3 and C4 have been shown to correlate strongly with active lupus, particularly with renal disease (31). A complete blood count, metabolic panel and urinalysis looking for involvement of target organs should also be obtained in the evaluation of a lupus patient. Further imaging and diagnostic studies are reserved for those patients for more severe complaints or disease state.

5. Pharmacotherapy and the Indications for Treatment of Lupus

The types of therapy that will be instituted for a lupus patient will depend on the degree of clinical presentation and severity of disease. Patients that have arthralgias and myalgias may be managed by non-steroidal anti-inflammatory agents. Because lupus patients have a propensity for drug-induced reactions, judicious use of medications and weighing of the risks and benefits of such medication is paramount. It is worth noting that cases of meningoencephalitis due to the use of ibuprofen in patients with autoimmune diseases, especially lupus, have been reported (32).

Because corticosteroids have an abrupt onset of action, their role is reserved primarily for patients requiring immediate therapy and/or control of their disease state. It is often employed as bridging therapy with a subsequent taper once disease modifying drugs have become effective in these patients. The actual mechanisms of action of corticosteroids are complex and go beyond the scope of this monograph. Suffice to note that the general consensus is that corticosteroids’ mechanism of action and side effects are due to the modulation of gene transcription (33). It is never appropriate for an abrupt termination of steroids, even with those patients on low steroid doses, as this may lead to an addisonian crisis due to the suppression of the adrenal glands. Long term use of steroids is discouraged due to their inherent side effects such as diabetes mellitus, osteoporosis, avascular necrosis, glaucoma and cushingoid features.

Disease modifying anti-rheumatic drugs (DMARDs) are used for maintenance and long-term therapy of lupus patients. The profiles of these drugs range from slow-acting anti-rheumatic drugs (SAARDs) such as the anti-malarials (hydroxychloroquine), to anti-metabolites (methotrexate), to more immunosuppressive drugs (cyclophosphamide, moftel mycophenolate). Lupus patients with rashes and non-erosive arthritis benefit from the use of hydroxychloroquine. Hydroxychloroquine is now considered the gold standard for the control of this disease due to its beneficial effects on the suppression of flares, controlling dyslipidemia as and prolongation of survival rates in lupus patients (34). Methotrexate is often used as a steroid-sparing agent and/or as a maintenance drug after induction therapy with cytotoxic agents (35). Lupus patients with more severe systemic states involving renal, neurologic and/or vasculitis are treated with cyclophospham ide, mofetil mycophenolate or azathioprine (36, 37). For the first time in over half a century, a FDA-approved biologic agent, belimumab (Benlysta), now available for SLE patients, is a human monoclonal antibody which functions as a B-lymphocyte stimulator (BlyS) - specific inhibitor, blocking the binding of soluble BlyS, a B-cell survival factor, to its receptors on B cells (1). It has long been speculated that B-cell modulation may be beneficial for lupus patients which has been the impetus for more investigational agents (38). Belimumab was studied and approved for adult patients with active, autoantibody-positive SLE receiving standard therapy such as NSAIDs, steroids, anti-malarials and immunosuppressives. It is administered as infusion therapy: IV 10 mg/kg at 2-week intervals for the first 3 doses and at 4-week intervals thereafter. The trial noted the adverse effects of increased infections, nausea, diarrhea, depression and even mortality more than placebo (39). One of the major criticisms of the BLISS study was the exclusion of lupus patients with kidney and CNS involvement from the trial. In addition, African American & Hispanic patients in the study were a small sample size and did not appear to respond to treatment with belimumab (Benlysta). The study lacked sufficient numbers to establish a definite conclusion in these particular patients. However, extension trials for this subpopulation of patients may help evaluate the safety and effectiveness of Benlysta.

6. Conclusions

Lupus is a chronic inflammatory autoimmune disease which can affect any organ system, but mainly involves the skin, joints, kidneys and nervous system. The diagnosis of lupus and the recognition of its various subtypes must be based on the proper constellation of clinical findings and laboratory evidence. Recognition of the pathogenic auto-antibodies and their targeted organs can help predict outcomes and treatment responses. Continual monitoring parameters include periodic follow-up and laboratory
testing to detect signs and symptoms of any new organ-system involvement. Overall treatment will depend on disease severity and organ involvement weighing the risks and benefits of each treatment modality.

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