Chapter 1

BILOGIC THERAPIES FOR SYSTEMIC LUPUS ERYTHEMATOSUS

Maria Mouyis1, MBBS MRCP, Coziana Ciurtin2,3, MBBS MSc PhD, and David Isenberg2,3,* MD FRCP FAMS

1Department of Rheumatology, Northwick Park Hospital, London, UK
2Department of Rheumatology, University College London Hospital NHS Foundation Trust, London, UK
3Centre for Rheumatology, Department of Medicine, University College London, London, UK

ABSTRACT

Advances in molecular biology have led to the development of biologic therapies. This is particularly relevant in systemic lupus erythematosus (SLE), which is a multisystem autoimmune rheumatic disease (ARD) associated with potentially life-threatening complications if not adequately treated. The availability of new biologic drugs has improved the prognosis of SLE in selected cases associated with

* Correspondence to: Prof. David A Isenberg, Academic Director of Rheumatology, Centre for Rheumatology, 424 The Rayne Institute, 5 University Street, University College London, London, WC1E 6JF, UK, email: d.isenberg@ucl.ac.uk
unsatisfactory response to conventional therapies. Over the last decade, there have been developments in the availability of biologic agents for SLE treatment based upon the advances in the understanding of the disease pathogenesis. Even if the evidence of biologic treatment efficacy in SLE is weaker than in other autoimmune rheumatic diseases, such as rheumatoid arthritis (RA), significant progress was made, as the first biologic treatment for use in SLE patients received approval in 2011. These new biologic therapies for SLE range from anti-CD20/CD22 (clusters of differentiation characteristic to B cells), to anti-B cell activating factors and anti-interferon alpha (IFNα). This chapter reviews the various biologic agents used in SLE, their mechanism of action and safety profile. The most common side effects to biologic treatments include infection, tuberculosis (TB) reactivation and allergic reactions. Less common side effects include development of lymphoma and anti-drug or autoimmune antibody formation. Despite their toxicity profile, biologic agents are gaining ground in clinical practice, due to the limited efficacy or increased toxicity of conventional disease modifying agents (DMARDs). Biologic therapies targeting B cells, such as rituximab, and B cell activation factors, such as belimumab, are currently used in the treatment of refractory SLE. Furthermore, aggressive treatment, including the use of biologic agents, reduces long-term complications associated with prolonged use of steroids in SLE, such as cardiovascular disease and osteoporosis. In the short term, the biologic agents are expensive when compared to traditional DMARDs; however there is evidence that their use is associated with long term benefits for patients with SLE, such as reduced hospital admission and disease complications, and improved patient outcomes. This chapter provides a summary of most biologic agents tested in SLE patients, considering their efficacy and safety profile, as well as the health implications associated with their use. We also take a brief look at newer agents currently investigated in clinical trials.

Keywords: systemic lupus erythematosus, biologic treatments, safety, efficacy, biosimilars.

INTRODUCTION

SLE is a complex ARD characterised by clinical manifestations that range from mild to severe. For many years the main treatments used were the traditional DMARDs, such as hydroxychlorquine, azathioprine, cyclophosphamide, methotrexate and mycophenolate mofetil (MMF). Unfortunately, these therapeutic agents have increased toxicity or, in some cases, limited efficacy in controlling the complex symptoms of this disease.
Despite the progress made in optimising the treatment of severe disease manifestations (e.g., lupus nephritis), the long-term prognosis of this disease has not changed dramatically in the last 30 years [1]. Patients with SLE are often treated for many years with prednisolone, which provides an additional array of complications, such as hypertension, glaucoma, steroid induced diabetes and osteoporosis. Although the treatment of SLE has, on the whole, improved lupus outcomes, less success was achieved in preventing or addressing the increased morbidity of this condition [2-4]. The availability of biologic agents is leading to a new treatment era for SLE patients, and there is hope for a better therapeutic management in the future.

**PATHOGENESIS**

A deeper understanding of the pathogenesis of SLE facilitated the discovery and development of many biologic agents targeting various molecules or receptors [5]. The correlation between different cellular and molecular players identified as key factors in lupus pathogenesis, and the available biologic therapies targeting the abnormalities associated with lupus, are illustrated in Figure 1. SLE is largely a B cell driven phenomenon with interplay between genetic, hormonal and environmental factors [6]. External triggers, such as ultraviolet (UV) radiation or Ebstein-Barr virus (EBV) infection, in conjunction with a maladaptive immune system and altered epigenetics are known to lead to the accumulation of apoptotic nuclear debris comprising anti-double stranded DNA (dsDNA) fragments and RNA antigens [7-9]. The debris is then processed by B cells, which function as antigen presenting cells (APC). This process then precipitates the formation of antibody production, and the release of pro-inflammatory cytokines such as interleukin (IL) 6, IL10, interferon γ (IFNγ), B lymphocytes stimulator/a proliferation-inducing ligand (BlyS/APRIL) and tumour necrosis factor alpha (TNFα). These cytokines promote auto B cell and auto T cell activation leading to a sustained inflammatory response [10].

T cells are important in the homeostatic control of the B cell responses. In SLE patients T cells are dysregulated. T1 helper cells are overexpressed and release significant amounts of IL12, IL18 and IFNγ. T2 helper cells are not overexpressed in comparison to T1 helper cells, but they secrete increased IL10 levels [11, 12]. There is also a decreased production in IL2 levels [13]. Regulatory T cells (Tregs), which are meant to dampen the pro-inflammatory T cell profile, are suppressed in SLE [14].
Legend: APRIL - *a proliferation-inducing ligand*; BAFF – *B cell activating factor*; BlyS – *B lymphocytes stimulator*; CD20 – *cluster of differentiation 20 marker* expressed from late pro-B cells through memory cells, but not on either early pro-B cells or plasma blasts and plasma cells; CD22 – *cluster of differentiation 22* found on the surface of mature B cells and to a lesser extent on some immature B cells; CD80 – *cluster of differentiation 80* is a protein found on activated B cells and monocytes that provides a costimulatory signal necessary for T cell activation and survival; IL – *interleukin*.

**Figure 1.** Schematic presentation of targeted therapy of SLE.

Furthermore there are changes in signalling subunits on the T cell receptors which lead to an increase production of co-stimulatory molecules such as cluster of differentiation 40 ligand (CD40L) [15]. The role of T cells in SLE is still being researched and new target biologic therapies are currently explored.

Another pathogenic process associated with lupus is the formation and deposition of immune complexes [16]. In SLE there is a failure of clearing immune complexes leading to immune complex deposition in organs such as kidneys or blood vessels, causing inflammation and tissue injury [17].
Every aspect of the immune abnormalities associated with lupus can be theoretically explored for therapeutic purposes. Progress has been achieved in testing different experimental biologic drugs. Recently, belimumab, an anti B cell activating factor (BAFF) therapy, became the first biologic treatment ever approved by Food and Drug Administration agency (FDA) for use in SLE patients, and the only treatment for lupus approved in over 50 years. It has also been approved for use in SLE patients in Europe, Middle East and Africa. Despite the impressive advances in molecular biology and drug technology, this complex autoimmune disease is still associated with increased morbidity and mortality. New treatment options are continuously explored to address the unmet need of a better quality of life and outcomes for patients (Figure 1).

**BILOGIC THERAPIES**

The biologic therapies, currently in use or under development for lupus target B cells, T cells, IFNα, IL6, and fusion proteins. This chapter explores the main biologic therapies available for the treatment of SLE patients, including the agents currently under investigation, detailing their mechanism of action, side effects and dosages. The level of evidence of efficacy and safety related to the efficacy of these biologic agents is summarised in Table 1.

**Advantages and Disadvantages**

The advantages of biologic therapies compared with some of the conventional therapies used for moderate-severe manifestations of SLE include potential efficacy in terms of disease control and relative safety in pregnancy, especially when compared to methotrexate, MMF and cyclophosphamide, which are contraindicated in pregnancy. The biggest disadvantage of both, biologic and non-biologic DMARDs, is that of immunosuppression, which is associated with increased risk of infection. Patients treated with biologics have to be screened for TB and hepatitis, as both infections can reactivate during the biologic therapy administration. A more significant concern, with most biologic agents, is the risk of progressive multifocal leukoencephalopathy (PML). This is not a drug specific phenomenon; although extremely rare, it is associated with significant mortality [18].

Another concern is that of the cost of biologic therapies. Biological agents are expensive as they are manufactured through advanced processes of
biotechnology and genetic engineering. However, the cost of treatment needs to be compared to the cost of having to manage all the challenging aspects of the disease. SLE patients in general utilise more health resources according to the severity of their disease and age at diagnosis, and patients with active lupus nephritis or regular flares even more so than a patient with mild or quiescent disease [19-21].

The cost of a patient having SLE includes direct and indirect costs [22]. The indirect costs are related to an individual’s quality of life and ability to work [23]. The cost of an individual’s loss of work on average in the US is was estimated at approximatively $16,345 per year in 2011 [19]. About 10 years ago, direct costs were assessed in a tri-nation study, which evaluated patients with SLE from several tertiary centres in Canada, US and UK respectively [24]. The direct cost of lupus treatment, including productivity over a 1 year period, was approximately $20,000 in the US [24]. There were also significant differences of the health system indirect costs in different countries [25, 26], and differences in the access of patients with lupus to the health care system across the world [27, 28]. A study from 2008 estimated that the annual (direct and productivity) cost for patients with lupus of employment age was $20,924 [23]. The cost of rituximab is approximately £4000 for two infusions in the UK, and there are data suggesting the cost-saving potential of rituximab in the management of lupus nephritis [29]. With the advent of biosimilars, the cost of treating SLE may become cheaper, but the efficacy of biosimilars (arguably having similar properties to the “original” biologic treatment) has yet to be proven [30, 31]. In the context of discussing the cost-effectiveness of biologic treatments it is worth mentioning that another advantage of the use of biologic agents is the intravenous (IV) administration of some of the available agents, which improves patient compliance and adherence to medication. New therapies, such as atacicept are already available as subcutaneous (SC) injections; therefore the above advantage might be lost for some of biologic therapeutic options. On the other hand, preserving the patient independence and enabling them to self-administer injectable treatments can contribute in itself to the reduction of additional health care costs. A study from 1991 by Petri et al. reported poorer adherence amongst Afro-Caribbean patients leading to more significant renal SLE and indirect increase in the healthcare costs [32]. Ultimately the cost of treatment needs to be balanced against the economic cost of disease for a realistic estimate of the cost-efficacy of different biologic therapeutic options.
Table 1. An overview of biologic therapy in SLE

<table>
<thead>
<tr>
<th>Biologic Agent</th>
<th>Trial</th>
<th>Mechanism of action</th>
<th>Side effects</th>
<th>Dosage</th>
<th>Organ specific indication</th>
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<tbody>
<tr>
<td><strong>B-cell depleting agents:</strong></td>
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<tr>
<td>Rituximab</td>
<td>Explorer Lunar</td>
<td>Anti-CD20 chimeric monoclonal antibody</td>
<td>Infusion reaction, Increased infection risk, PML, Lymphoma</td>
<td>1 g IV twice, 2 weeks apart 375 mg/m² IV every 4 weeks 750 mg cyclophosphamide may be given with the first infusion to increase B cell depletion effect</td>
<td>SLE nephritis Non-renal SLE (despite negative RCTs results)</td>
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<tr>
<td>Ofatumumab</td>
<td>RA trials, Case reports in SLE</td>
<td>Fully human monoclonal antibody against membrane proximal epitope on CD20 molecule</td>
<td>Infusion reactions: Urticaria, Rash, Rhinitis, Nausea, URTI, Headaches, Fatigue, Flushing</td>
<td>300, 700, 1000 mg IV every 2 weeks for 24 weeks.</td>
<td>Arthritis (RA) ?SLE</td>
</tr>
<tr>
<td>Ocrelizumab</td>
<td>Phase III trials in SLE</td>
<td>Fully human monoclonal antibody against CD20</td>
<td>Severe infections (increased in patients treated with MMF)</td>
<td>400 mg or 1,000 mg ocrelizumab, given as an IV infusion on days 1 and 15, followed by a single infusion at week 16 and every 16 weeks thereafter</td>
<td>SLE nephritis</td>
</tr>
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<td>Biologic Agent</td>
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<td><strong>Anti-B-cell activating factors</strong></td>
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<td>Epratuzumab</td>
<td>EMBODY-1 EMBODY-2</td>
<td>IgG1 monoclonal antibody against the CD22 molecule</td>
<td>Infusion reaction, URTI’s Fever Headache Nausea and dizziness</td>
<td>600 mg IV every week for four weeks or 1,200 mg IV every two weeks for four weeks</td>
<td>Neuro-psychiatric, mucocutaneous and musculoskeletal SLE manifestations.</td>
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<tr>
<td>Belilumab</td>
<td>BLISS 52 BLISS 76</td>
<td>Human monoclonal immunoglobulin (IgG1y) against BAFF/BLySS.</td>
<td>Nausea Diarrhoea Headaches URTI Fever Cystitis Infusion reaction</td>
<td>10 mg/kg IV every 2 weeks x 3 and then once every 4 weeks</td>
<td>Non renal SLE Non cerebral SLE</td>
</tr>
<tr>
<td>Tabalumab</td>
<td>IILUMINATE</td>
<td>Human monoclonal antibody against BAFF</td>
<td>URTI UTI Injection site reactions Myocardial infarct Discitis Osteomyelitis Breast cancer Cerebrovascular accident Pulmonary fibrosis</td>
<td>240 mg SC loading dose, followed by 120 mg SC every 2 or 4 weeks</td>
<td>Non-renal SLE Non-cerebral SLE</td>
</tr>
<tr>
<td>Blisibimod</td>
<td>Phase II PEARL-SC</td>
<td>Fusion protein, selective antagonist of BAFF</td>
<td>Injection site reactions</td>
<td>200 mg SC weekly</td>
<td>Non-renal SLE Non-cerebral SLE</td>
</tr>
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<tr>
<td><strong>Atacicept</strong></td>
<td>Phase II APRIL-SLE (terminated)</td>
<td>TACI-Ig fusion protein that inhibits BLyS and APRIL</td>
<td>LRTI/URTI Injection site reaction Fever Arthralgia Sinusitis Headache Fatigue Rhinitis Dizziness Depression</td>
<td>75 mg or 150 mg SC weekly</td>
<td>SLE nephritis-study terminated because of side-effects.</td>
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<td>Phase II ADDRESS-II (ongoing)</td>
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<td><strong>Anti-IFN alpha</strong></td>
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<td>Sifalimumab</td>
<td>Phase III trial</td>
<td>IFNα monoclonal antibody</td>
<td>Infusion reaction Fatigue URTI UTI Sinusitis Dizziness Arthralgia Headache Lymphopenia Anaemia</td>
<td>0.3, 1.0, 3.0, or 10.0 mg/kg IV- still undergoing trials</td>
<td>SLE nephritis</td>
</tr>
<tr>
<td>Rontalizumab</td>
<td>Phase II trial</td>
<td>A recombinant humanized monoclonal antibody to IFNα</td>
<td>Viral infections Reactivation of Herpes infection Sinusitis Bronchitis</td>
<td>750 mg IV 4 weekly 300 mg SC 2 weekly 0.3-10 mg/kg</td>
<td>Moderate-severe SLE</td>
</tr>
</tbody>
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Table 1. (Continued)

<table>
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<tr>
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</tr>
</thead>
<tbody>
<tr>
<td>Anifrolumab</td>
<td>Phase II trial</td>
<td>A type I IFN receptor antagonist</td>
<td>Influenza, Herpes-Zoster</td>
<td>300 mg IV monthly, 1000 mg IV monthly</td>
<td>Moderate-severe SLE</td>
</tr>
</tbody>
</table>

**Blockade of T cell activation**

| Abatacept | Phase III trials | Human IgG1 heavy chain fused with CTLA4 that blocks T cell activation by B cells. | Nausea, Headache, Infusion reaction, Fever, Hypertension, Back pain, Infections | 10-30 mg/kg IV | Discoid SLE, Arthritis, Serositis |

**IL6 inhibition**

| Tocilizumab | RA trials, Phase 1 SLE trials | Monoclonal IgG1 antibody to IL6 receptor | URTI, GI infections, TB, GI perforation, Non-melanoma skin tumours, Malignancies, Abnormal LFT, High cholesterol levels, Suppression of CRP | 4 or 8 mg/kg IV | SLE nephritis, SLE with moderate activity |

Legend: CRP - C reactive protein; CTLA4 - cytotoxic T-lymphocyte-associated protein 4; GI - gastro-intestinal; IFR - individual funding request form, IFN - interferon; Ig - immunoglobulin; IL - interleukin; IV - intravenous; LFT - liver function tests; LRTI - lower respiratory tract infection; MMF - mycophenolate mofetil; PML - progressive multifocal leukoencephalopathy; RA - rheumatoid arthritis; RCT - randomized controlled trial; SC - subcutaneous, SLE - systemic lupus erythematosus; TB - tuberculosis; URTI - upper respiratory tract infection; UTI - urinary tract infection.
The use of biologic agents in different countries is highly variable. In countries with poorer economic status, the access to expensive treatment options is limited and not always fairly distributed in the population. Mortality in lupus was associated with lower education, shorter duration of follow up and poor medical coverage in a large study from South American countries [33].

Belimumab is licensed for use to treat SLE in the US, Europe, but in the UK, the funding of the treatment is based on an individual funding request (IFR). The disparity between the outcome of patients with lupus nephritis with private vs. public health insurance in US is well recognised [27, 34].

Belimumab has not been approved by NICE (National Institute of Clinical Excellence) who do not consider it cost-effective as assessed by QALY (quality-adjusted life year) calculations; although real life experience shows some clinical benefits in patients with skin and joint manifestations [35]. Arguably QALY, which aims to appreciate the disease burden by taking into consideration both, the quality and duration of life, is a flawed measurement tool in the healthcare system [36]. An ideal system treatment for SLE would ensure an increased good quality life span at the lowest possible health care cost. This challenge is compounded by the dilemma of using biosimilars, which supposedly have similar efficacy as the biologic agents and the advantage of lower cost. However, data regarding the efficacy of biosimilars in head-to-head clinical trials aiming to compare them with the original biologic agent are missing for the majority of available options.

Table 1 summarises the most studied biologic treatments in patients with lupus, with particular emphasis on their efficacy for certain SLE manifestations, dosage and toxicity profile.

### Biologic Therapies Affecting B Cells

1. Anti-CD20 - rituximab, ofatumumab, ocrelizumab, veltuzumab
2. Anti-CD22 - epratuzumab

#### Rituximab

**Mechanism of Action and Dosage**

Rituximab is a chimeric/humanised monoclonal antibody against CD20. It was the first biologic to be used in the treatment of SLE. The drug depletes CD20 B lymphocytes. The CD20 antigen is found on pre-B cells which would then form mature B cells [37]. Depletion of the B cell therefore reduces cell apoptosis and complement activation [38, 39]. Although the function of CD20 is unknown, it is considered that it may play a role in Ca\(^{2+}\) influx across plasma
membranes, maintaining intracellular Ca\(^{2+}\) concentration and allowing differentiation and activation of B cells.

The fragment antigen binding (Fab) domain of rituximab binds to the CD20 antigen on B lymphocytes, and the fragment crystallisable (Fc) domain recruits immune effector functions to mediate B cell lysis in vitro; in addition rituximab increases the expression of major histocompatibility complex II (MHC II), and lymphocyte function-associated antigens 1 and 3 (LFA-1, LFA-3), triggers shedding of CD23, down-regulates B cell receptor and induces the apoptosis of CD20 B cells [40].

The standard dose currently recommended for the treatment of SLE is 1 g of IV rituximab given 2 weeks apart. Each dose is preceded by premedication with methylprednisolone, antihistamine and paracetamol to reduce the risk of infusion reactions. Cyclophosphamide may also be given pre-first dose to increase the efficacy of rituximab and enhance B cell depletion [41].

**Efficacy and Side Effect Profile**

Rituximab is considered effective in treating refractory SLE, although two large trials, LUNAR (clinical trial of patients with lupus nephritis) and EXPLORER (clinical trial which included non-renal patients) did not meet their primary endpoints. The reason for this is likely related to the concomitant use of high dose of steroids and concomitant conventional immunosuppressant therapy [42, 43]. However, some significant clinical and serological results were noted (e.g., significant reduction in proteinuria, lower steroid requirement, etc.). Despite the failure of these clinical trials, numerous case-reports and studies using rituximab off-label in lupus patients have shown a 70-90% response rate [43-46]. The role of Rituximab as a steroid sparing agent has also become evident. Condon et al. reported results on 50 lupus nephritis patients treated with 1 g of rituximab 2 weeks apart and 500 mg of IV methylprednisolone [47]. No oral steroids were given and MMF was used as maintenance therapy. 70% (n = 36) of the patients achieved remission in a mean time of 36 weeks. The conclusion of the researchers was that oral steroids can be avoided (only 2/50 patients required them in a 2 year follow up period), and B cell depletion might be considered early in the treatment of lupus nephritis and other systemic manifestations of SLE. Rituximab minimized the need for additional steroids and was effective in ensuring a better long term control of the disease. An earlier, albeit smaller study of eight newly diagnosed rituximab treated patients (each case carefully matched to three conservatively treated patients) reported similar steroid sparing capacity of rituximab in non-renal lupus patients [48]. An international trial comparing rituximab and MMF in lupus nephritis patients
treated with minimal dose of steroids vs. standard therapy with high doses of steroids and MMF (the RITUXILUP study) is currently undergoing and should provide more definitive answers regarding the exciting possibility to treat patients with lupus with low doses of steroids and rituximab.

Side-effects of rituximab include infusion reactions as documented in the initial oncology studies [49]. The most common side-effects are fever, bronchospasm, rash and hypotension, which usually settle on stopping the infusion. Patients are usually followed up carefully post-rituximab treatment to monitor for the development of common bacterial and viral infections, or other infections, such as TB and hepatitis B or C. Based on the experience acquired from treating RA patients with rituximab, this biologic agent is considered reasonably safe in the context of history of TB [50, 51]. There are reports of both reactivation of hepatitis B following treatment with rituximab [52], as well as effective treatment of vasculitis associated with hepatitis C with rituximab [53, 54]. Human anti chimeric antibodies have also been reported [55], and side-effects to medication are considered to be linked to the immunogenicity of these antibodies [56]. The effect of B cell depletion lasts for 6-12 months in about 75% of cases, and the response to therapy is variable [38, 57, 58], and, as shown recently, the longer the duration of B cell depletion, the better the outcome [59]. The process of B cell repopulation following B cell depletion therapy with rituximab in lupus is still not fully understood [60]. For safety reasons, it is recommended to check immunoglobulin (Ig) levels and CD19+ B cell count every 2 months until B cells normalise, as accumulated doses of rituximab may also cause hypogammaglobulinaemia which may be linked to an increased the risk of infection [61, 62].

The expert consensus is that rituximab is a safe and effective treatment option for patients with refractory renal and non-renal lupus and can reduce significantly the steroid burden [48, 63-65], despite the negative trial results.

**Ofatumumab**

*Mechanism of Action and Dosage*

Ofatumumab is a fully human monoclonal antibody anti-CD20 [66]. The treatment is licensed for use in patients with chronic lymphocytic leukaemia based on its proven efficacy in treating refractory cases [67, 68]. Most studies assessing dosing regimens in patients with rheumatic conditions have been done in RA patients, in whom the treatment was also associated with clinical benefit [69, 70]. The current recommended dose is 700 mg IV every 2 weeks. Similarly
with other biologic agents, premedication is administered beforehand, to minimise the risk of infusion reactions [71].

**Efficacy and Side Effect Profile**

The safety profile and efficacy of ofatumumab has also been established in a few RA clinical trials [69, 71]. The drug has been proved effective as B cell depletion agents as was associated with significant clinical improvement of symptoms of arthritis at different doses [71]. The pharmacokinetic of the medication was found similar in chronic lymphocytic leukaemia and RA [72]. A recent case report has suggested some benefit in the treatment of SLE [73].

Although safety profiles were established in phase 1 and II trials, the most comprehensive information related to the drug’s safety profile was provided by a phase III clinical trial in RA [69]. The most common side-effects were rash (21%) and urticaria (12%), which mostly occurred on the day of first infusion and they declined significantly with the second course. Most adverse events were of mild or moderate intensity (see Table 1).

**Ocrelizumab**

**Mechanism of Action and Dosage**

Ocrelizumab is another fully human monoclonal antibody against CD20, developed for RA, SLE, and B cell derived malignancies [74, 75], which was tested for efficacy in patients with lupus nephritis [76]. The doses used in the largest phase III clinical trial were 400 mg or 1,000 mg ocrelizumab, given as an IV infusion on days 1 and 15, followed by a single infusion at week 16 and every 16 weeks thereafter [76].

**Efficacy and Side Effect Profile**

Despite reaching an overall response rate of 66-67% in the ocrelizumab treatment arm, the difference in response vs. standard of care treatment did not reach statistical significance [76]. The study was terminated earlier as there was an infection-related safety signal in relation with increased risk of opportunistic and fatal infections in the ocrelizumab treatment groups [77]. The proportion of patients experiencing serious infections was twice as high in patients who received concomitant treatment with MMF (32% vs. 16% in the placebo arm), and it was increased in Asian patients [76].

Although ocrelizumab has been unsuccessful in one clinical trial in SLE, this negative result may be attributed to the use of concomitant MMF. It has also been trialed in relapsing and remitting multiple sclerosis with only 3 of 55
patients experiencing significant adverse events. The treatment was associated with a reduction in neurological lesions on magnetic resonance imaging (MRI) [78]. It remains possible that this drug can still be used in SLE pending additional trials.

**Veltuzumab**

*Mechanism of Action and Dosage*

Veltuzumab is a second generation humanised anti CD20 monoclonal antibody with a SC formulation, developed for the treatment of refractory pemphigus vulgaris and hematologic malignancies [79, 80]. A case report showed improved serological outcome with treatment with veltuzumab in a lupus patient who developed anti-drug antibodies (HACA) to rituximab [81]. The role of veltuzumab in SLE has yet to be clearly determined but may be used where rituximab is ineffective or there is “resistance” as seen in patients with non-Hodgkin lymphoma (NHL) [82].

**Bispecific Monoclonal Antibodies**

Recent laboratory studies suggested that the use of bispecific antibodies (anti CD22/CD20) might be effective in the treatment of lupus, as they were associated with increased trogocytosis. Trogocytosis is a process whereby lymphocytes conjugated with APC extract surface molecules from APCs and express them on their own surface in vitro, resulting in decreased levels of B cell surface markers associated with considerably less B cell depletion and therefore less risk of severe immunosuppression [83].

**Epratuzumab**

*Mechanism of Action and Dosage*

CD22 is a B cell transmembrane glycoprotein that is found on mature B cells. Epratuzumab is an IgG1 monoclonal antibody against the CD22 molecule, which inhibits B cell receptor activation and leads to subsequent B cell apoptosis. The ALLEVIATE-1 and ALLEVIATE-2 randomised clinical trials (RCTs) proved that 360 mg/m² IV dose was more effective than the 720 mg/m² IV dose as a steroid-sparing medication in patients with severe lupus. The responses correlated with improvements in health-related quality of life [84]. The EMBLEM study was an early phase II B study performed to determine the
effective dose regimen in patients with moderate-severe lupus, which showed that patients receiving a total of 2400 mg had significant improvement in the disease activity [85].

The phase III studies, EMBODY 1 and 2, enrolled SLE patients who received placebo or treatment with 2400 mg of epratuzumab over four 12-week treatment cycles, administered as 600 mg every week for four weeks or 1,200 mg every two weeks for four weeks. The primary endpoint of both studies (which was defined as the percentage of patients meeting treatment response criteria at week 48 according to the British Isles Lupus Assessment Group (BILAG) - based Combined Lupus Assessment - BICLA) was not met and the research program was discontinued. BICLA, like the SRI (SLE responder index) system, requires patients to meet response criteria across three assessment tools: the BILAG-2004 index, SLEDAI index (SLE- disease activity index) and a physician’s global assessment (PGA).

Epratuzumab is considered to reduce on average only 35% of circulating B cells in patients, and has minimal antibody and complement-dependent cellular cytotoxicity (when evaluated in vitro), and it was hypothesized that its therapeutic activity may not result completely from B cell depletion [83]. However, this was not translated in clinical benefits.

Side Effect Profile

Epratuzumab has been used in the treatment of both SLE and Sjögren’s syndrome [85-87]. The drug has been proven to reduce BILAG scores as well as neuropsychiatric (62.5%), muco-cutaneous (21.9%) and musculoskeletal symptoms (32%) compared to placebo [85].

The common side effects included infusion reaction, upper respiratory tract infections (URTI’s), fever, headache, nausea and dizziness. A small proportion of patients developed human antidrug antibodies (HAHA). In contrast to rituximab, no severe decrease in the Ig levels was noted and this may be in relation to a partial depletion of B cells.

Anti B Cell Activating Factors

1. Belimumab
2. Tabalumab
3. Atacicept
Belimumab

Mechanism of Action and Dosage
This is a monoclonal humanised Ig which binds to the BLyS protein. It is a transmembrane protein expressed by T cells, dendritic cells and neutrophils. The BLyS protein binds to 3 receptors on the B cell surface: the B cell maturation antigen (BMCA), BAFF-receptor (BR3, BAFF-R) and a transmembrane activator and calcium modulating ligand interactor (TACI); via these receptors it has a role is in B cell differentiation, Ig production and levels of disease activity [88]. Patients with SLE have high levels of BLyS, therefore binding of the BLyS protein leads to decrease B cell activation, maturation and antibody production. The dose used in lupus clinical trials was 10 mg/kg IV every 2 weeks for 6 weeks and thereafter monthly.

Efficacy and Side Effect Profile
Belimumab is considered effective in patient with non-renal and non-cerebral SLE. FDA has approved it for mild to moderate SLE for those with skin and joint disease. It is not approved for active renal lupus or cerebral lupus. BLISS 52 and BLISS 76 studies have shown efficacy at 52 and 76 weeks [89]. BLISS 52 was carried out in countries from Central and Eastern Europe, Asia-Pacific, and Latin America. At 52 weeks, there was a 58% patients who had been treated with 10 mg/kg of belimumab met the primary outcome, which was the SRI response rate (P = 0.017). SRI comprises criteria from three different internationally validated indices, SELENA-SLEDAI, PGA and BILAG 2004. BLISS 76 was a clinical trial which included patients from North America, and Western and Central Europe. Similarly, at 52 weeks 43% of patients treated with 10 mg/kg of belimumab met a similar primary outcome (p < 0.02). The 52 week response was not maintained at 76 weeks. The results of these large clinical trials suggested that improvement of disease control is more likely in patients with low C3 and high dsDNA levels [90]. A pooled post-hoc analysis of the combined phase III studies suggested a possible benefit in lupus nephritis as well [91]. Belimumab was associated with improvement of disease activity, reduced flares, decrease in dsDNA levels and low rate of side effects and it is currently licensed for use in non-renal lupus patients. The one drawback of belimumab is the delay onset of action and therefore not ideal for an acute flare treatment. It can take up to 6 months to achieve 70% B cell depletion. Common side effects included nausea, diarrhoea, headaches and URTIs, but their frequency was low. The rates of patients experiencing adverse events, as
assessed from pooled data from one phase II and two phase III RCTs were 16.6%, 19.5%, 13.5%, and 18.0% with placebo, and belimumab 1, 4, and 10 mg/kg, respectively [92]. There was also evidence of low rate of serious infusion reactions (including hypersensitivity reactions) occurring at a lower frequency that 1% in both placebo and active medication patient groups [92]. The side effect profile is not dose dependent. There is one case report of a severe delayed anaphylactic reaction which was fatal [92].

**Tabalumab**

*Mechanism of Action and Dosage*

BLyS, also known as BAFF, belongs to the TNF family. It is expressed by multiple cells including macrophages, monocytes, neutrophils and dendritic cells. BLys/BAFF binds to 3 receptors called BR3, thus affecting B cell lineage. TACI receptor is found on T cells and marginal zone B cells. BCMA (B cell maturation antigen) receptor is found on plasma cells. Activation of these receptors leads to auto-antibody production, increase in B cells and possibly potentiate malignancies [93].

In contrast to belimumab, tabalumab is an anti BAFF monoclonal antibody, which targets both membrane bound and soluble BAFF, currently used in RA [94]. The ILLUMINATE-1 study, a phase III RCT of tabalumab in patients with SLE, used the following dose regimen: 240 mg SC loading dose, followed by 120 mg every 2 weeks or 4 weeks in combination with traditional SLE treatments. The primary endpoint was the proportion of patients achieving an SLE Responder Index 5 (SRI-5) response at week 52. The SRI-5 is a composite endpoint defined as ≥5 point improvement (reduction) in SELENA-SLEDAI score, no new BILAG 2004 index score of A or no more than one new BILAG B score, and no worsening (increase ≥0.3 points from baseline) in PGA [95]. The study did not meet the primary endpoint. Similarly, no difference in the secondary outcomes (which included time to first severe SLE flare on the SELENA-SLEDAI Flare Index, proportion of patients with reduction in corticosteroid dose by ≥25% to ≤7.5 mg/day prednisone (or equivalent) for ≥3 consecutive months from weeks 24 through 52, and change from baseline on the Brief Fatigue Inventory at week 52), was found between the active and placebo groups. However, in a sensitivity analysis which did not exclude the patients who decreased antimalarial, steroid or immunosuppressant therapy, SRI-5 response was achieved with tabalumab 120 mg every 4 weeks (37.0% vs 29.8% placebo; p = 0.021), suggesting a potential for this biologic agent to be effective in selected categories of lupus patients [95]. Furthermore, the ILLUMINATE-2
study, which was set up in a broadly similar way to ILLUMINATE-1, but not excluding these patients whose concomitant medications were reduced, did not meet its primary endpoint. Unfortunately, despite the encouraging results, Eli-Lilly have terminated the tabalumab program in lupus.

**Efficacy and Side Effect Profile**

The side-effects noted from the initial RA trials include infections and injection site reactions [96]. The most common infections included URTI and urinary tract infections (UTIs). Severe adverse events were also noted, such as myocardial infarct, discitis, osteomyelitis, breast cancer, cerebrovascular accident and pulmonary fibrosis [96, 97]. The treatment with tabalumab was associated with slight increase in the incidence of depression and suicidality compared to placebo in the ILLUMINATE-1 study, but these side-effects were uncommon [98]. The incidence of serious infections and severe infections were similar in the tabalumab and placebo groups in SLE patients [95].

**Atacicept**

**Mechanism of Action and Dosage**

This novel agent inhibits both BLyS and APRIL in B cells, affecting B cells ranging from immature to mature. It is a TACI-Ig fusion receptor protein [99]. As described above, by inhibiting BLyS and APRIL it causes a reduction in B cell proliferation, IFNγ and Ig production [100]. The doses used in the phase II/III RCT in lupus were either 75 mg or 150 mg atacicept SC biweekly for 4 weeks and then weekly versus placebo [101].

**Efficacy and Side Effect Profile**

In the APRIL-SLE phase II RCT, the 150 mg atacicept arm was terminated early due to two fatal infections [101]. Despite this, a post-hoc analysis of atacicept 150 mg has shown that this dose regimen reduced the incidence of flares and time to first flare compared to placebo (flare rate 37% vs. 54%, odds ratio - OR = 1.15 (0.73-1.8), and time to flare HR = 0.56, P = 0.009) [102]. There was no difference between atacicept 75 mg and placebo. The clinical response was accompanied by decrease in B cells, Ig levels and increase in complement levels. The 75 mg arm failed to meet the primary endpoint, defined as a significant decrease in the proportion of patients experiencing at least one flare of BILAG A or B [101].

The main safety concern regarding atacicept is that of potential increased incidence of hypogammaglobulinaemia and therefore, risk of infection. A study
in patients with lupus nephritis was terminated after the enrolment of only 6 patients because of the severe decreased in the level of Ig [103]. A closer look at the concomitant medication, showed that these patients’ hypogammaglobulinaemia developed when the patients were given MMF before they were treated with atacicept [104]. It was hypothesised that targeting APRIL in autoimmune disease might be associated with significant risk of toxicity [105]. Further phase II/III clinical trials of atacicept in lupus are currently undergoing (ADDRESS II) and should be able to provide additional information about the safety profile of this biologic agent. Apart from the two deaths encountered in the 150 mg atacicept group in the APRIL-SLE trial, the proportion of the serious infections was not statistically significantly different between the 75 mg atacicept arm when compared to placebo [101]. Furthermore, the death and injection rates were similar to those reported in the belimumab studies. The most common infections encountered included haemophilus influenzae pneumonia, legionella pneumonia and bacillus bacteraemia. Preclinical studies showed an increase in liver transaminases [101].

**Blisibimod**

*Mechanism of Action and Dosage*

Blisibimod is a fusion protein consisting of four BAFF binding domains fused to the Fc region of a human antibody, which acts as a selective antagonist of BAFF. Blisibimod selectively inhibits both soluble and membrane-bound BAFF.

*Efficacy and Side Effect Profile*

The efficacy and safety of blisibimod in subjects with SLE was investigated in a phase II RCT, PEARL-SC study, which found that the highest tested dose of 200 mg blisibimod administered SC once weekly was associated with increased SRI-5 response rates, but without reaching statistical significance when compared with placebo. The treatment was more effective in patients with SELENA-SLEDAI improvement of ≥8, and in a subgroup of patients with severe disease (SELENA-SLEDAI ≥10) [106].

Blisibimod was associated with a decrease in the number of naïve B cells (24-76%) and a transient relative increase in the memory B cell compartment in the phase 1 studies [107]. It was also associated with significant decrease of dsDNA, increase in the complement C3 and C4, and reductions in serum B cell levels in the PEARL-SC study [106]. Blisibimod is currently being tested in a
phase III study for SLE, CHABLIS-SC1, and a phase II study, BRIGHT-SC, for IgA nephropathy.

The treatment with blisibimod was safe, as the incidence of serious side-effects was similar to the placebo arm. Injection site reactions were reported more frequently with blisibimod compared with placebo, but they were mild (erythema) [106]. Taking into consideration this treatment’s serological benefits in SLE patients and acceptable safety profile, the results of the phase III study are awaited with interest as if proven more effective than belimumab, the treatment has a good chance to be the next licensed biologic treatment for lupus.

*Anti-Interferon Alpha (IFNα)*

1. Sifalimumab
2. Rontalizumab
3. Anifrolumab

**Mechanism of Action**

Sifalimumab and rontalizumab are anti-IFNα monoclonal antibodies. An increase in BAFF occurs via signalling of INFα. The signalling pathway is activated by the stimulation of the IFN-1 receptor. In SLE pathogenesis there is activation of type 1 IFN, which is associated with lupus nephritis [108, 109]. Neutralisation of IFNα will lead to a reduction of inflammation by a reduction in BAFF levels, mature B cells, antibody production and T cell activation [110, 111].

**Sifalimumab**

**Dosage**

An optimal dose is yet to be confirmed as trials are currently underway. The phase I RCT in lupus used the following doses: 0.3, 1, 3, 10 or 30 mg/kg as a single IV administration [112]. The results of a promising phase II RCT of Sifalimumab in SLE were presented at the 2014 American College of Rheumatology (ACR) annual meeting [113].

The patients were randomised to receive monthly IV doses of sifalimumab at 200, 600, or 1200 mg or placebo for 1 year based on their disease activity, IFN signatures and geographic region. The primary endpoint, defined as the percentage of patients achieving an SRI at day 365, was achieved in all the treatment active arms (sifalimumab 200, 600, and 1200 mg doses were associated with 58.3, 56.5, 59.8%, SRI response respectively, compared to
45.4% in the placebo group). Surprisingly, there were no significant changes of the dsDNA or complement levels despite the good response to treatment [113].

**Efficacy and Side Effect Profile**

The results from the early phase clinical trials showed a reduction in SLE disease activity [108, 113], and phase 3 trials are currently underway. Sifalimumab is considered safe although there have been reports of increase incidence of *herpes zoster* infection. Other side effects typically include infusion reactions, nausea, URTIs, UTIs, headache and arthralgia [108, 114].

**Rontalizumab**

**Mechanism of Action and Dosage**

Rontalizumab has a similar mechanism of action as sifalimumab [109]. In an early phase, dose-escalation study, patients were enrolled into dose groups ranging from 0.3 to 10 mg/kg, administered via IV or SC routes [115].

**Efficacy and Side Effect Profile**

A recent phase II studies with rontalizumab in lupus did not meet the criteria for efficacy, which were reduction in disease activity as assessed by the BILAG and SRI [116]. A phase I study showed a dose dependent decrease in the level of IFNα, but no decrease in levels of dsDNA. The side effect profile was deemed similar to placebo although an increase in viral infections was noted [115, 116]

**Anifrolumab**

**Mechanism of Action and Dosage**

Anifrolumab is a type I IFN receptor antagonist [117], which was recently tested in a phase II RCT in patients with SLE using two dose regimens: 300 and 1000 mg IV anifrolumab, monthly administration (ACR abstract data, Merrill et al., 2015 in press).

**Efficacy and Side Effect Profile**

The primary endpoint of this phase II RCT of anifrolumab in patients with moderate to severe SLE was a composite SRI response at day 169 with sustained reduction of the steroid dose (<10 mg/day dose maintained between days 85 and 169. Both treatment regimens (300 and 1000 mg anifrolumab) improved patients’ outcomes and reached the primary endpoint (34.3% and 28.8% respectively, vs. 17.6% placebo). Steroid dose reduction (<7.5 mg daily) at day
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365 was achieved by 26.6% patients in the placebo group vs. 56.4% in the 300 mg anifrolumab group (p = 0.001) and 31.7% in the 1000 mg anifrolumab arm (p = 0.59).

**Efficacy and Side Effect Profile**

The treatment with anifrolumab was associated with similar serious adverse events than placebo. A higher incidence of influenza (most unconfirmed) and a dose-dependent increase in herpes zoster were reported in the anifrolumab treatment arms compared to placebo.

The results from previous clinical trials with monoclonal antibodies to IFN have shown encouraging results and further trials are ongoing.

**Blockade of T Cell Activation**

**Abatacept**

**Mechanism of Action and Dosage**

This is a fusion protein which interferes with the co stimulatory interactions between B and T lymphocytes. Activated T cells express cytotoxic T-lymphocyte-associated protein 4 (CTLA4) which interacts with co-stimulatory receptor B7-1 (CD80). Abatacept is a combination of human IgG (Fc portion) and CTLA-4. It therefore blocks stimulation of B cells leading to a reduction in antibody formation and immune response [118]. The doses used in lupus trials range from 10-30 mg/kg [119-123]

**Efficacy and Side Effect Profile**

Initial murine studies showed improvement in lupus nephritis, proteinuria and autoantibody titres but this has not yet translated into human studies. Phase II/III trials in lupus nephritis did not meet outcome measures, although when the same data were analysed using different criteria (LUNAR trial response criteria) there was a 20% response rate in the abatacept arm compared to placebo [119, 121].

The side effect profile is comparable to other biologics and is detailed in Table 1.

**IL6 Blockage**

1. Tocilizumab
2. Sirukumab
**Tocilizumab**

**Mechanism of Action and Dosage**

In mouse models exogenous IL-6 has been shown to increase autoantibody production and progression of lupus nephritis. By blocking IL-6 there is a decrease in antibody formation, proteinuria and mortality. IL-6 is released by intrinsic kidney cells and causes mesangial cell proliferation, activation of T and B cells and autoantibody secretion [124]. High IL-6 levels are associated with SLE disease activity as well as dsDNA levels. By blocking IL-6 there is a decrease in inflammation, B cell differentiation and autoantibody production [124]. Binding of IL-6 to the receptor is prevented by tocilizumab, a fully humanised monoclonal antibody. It can bind to membrane bound or soluble IL-6R [125]. The starting dose is 4 mg/kg monthly and this can be increased to 8 mg/kg monthly, pending clinical response [124].

**Efficacy and Side Effect Profile**

Although a well-established treatment in RA and systemic juvenile idiopathic arthritis (JIA) [126], its role in SLE is yet to be established. A phase I clinical trial in SLE proved the safety and efficacy of tocilizumab in lupus patients [124].

The side effect profile is thought to be less severe than with other biologic agents. URTIs and gastrointestinal infections are most common. The treatment with tocilizumab is associated with suppression of C-reactive protein (CRP), haematological abnormalities, non-melanoma skin tumours, and malignancies. Gastro-intestinal perforation has been reported in phase III trials in RA. Liver dysfunction and increased levels of LDL and total cholesterol has also been reported in clinical trials in RA [127, 128]. In the only clinical trial in lupus, the treatment was associated with decreased neutrophil levels, but without major impact in increasing the risk of infections [124].

**Sirukumab**

Sirukumab is a humanised monoclonal antibody against IL-6, similar to tocilizumab [129]. The results of a proof of concept study were reported at the ACR meeting in 2014 [130, 131]. Preliminary data suggested some improvement of the patient-outcome measures and transient improvement in
clinical parameters [132]. The treatment with sirukumab was associated with a dose-dependent decrease in absolute neutrophil count and platelet count [133].

**FUTURE TREATMENT OPTIONS**

**Anti-Complement Therapies: Eculizumab**

Complement activation is strongly involved in the pathogenesis of SLE. The function of complement is to help clear immune complexes and a deficiency in this leads to the development of SLE. Eculizumab has been developed to inhibit terminal complement activation and maintain early complement function [134]. It is a monoclonal antibody against C5. By blocking C5 it prevents the formation of C5a and C5b and the formation of the terminal membrane attack complex [135]. This drug is in phase 1 trials and limited data suggest a delay in the onset of proteinuria and improved outcomes in patients with hemolytic-uremic syndrome after renal transplantation [136]. A recent case report also suggested clinical benefit in treating severe lupus nephritis in a paediatric patient [137].

**CONCLUSION**

The various clinical and laboratory abnormalities associated with SLE need tailored therapeutic interventions. Despite the large number of biologic treatments with potential efficacy for controlling different aspects of lupus disease, it is worth mentioning that only one biologic treatment, belimumab, was proven effective in large phase III clinical trials leading to the licensing of a new therapy for lupus. The strict inclusion criteria used in clinical trials suggest that even if shown effective, these treatments might only be useful for selected categories of SLE patients and the generalisation of the results from clinical trials is not indicated. However, despite the lack of efficacy in clinical trials, rituximab is widely believed to be effective in treating refractory SLE and it is currently widely used off license. Future research should lead to reconciliation between the clinical trial results and clinician expertise related to the use of biologic treatments for the benefit of SLE patients.
ACKNOWLEDGMENTS

The authors would like to thank to Dr. Marwan Bukhari, Consultant Rheumatologist, Royal Lancaster Infirmary, Lancaster, UK (email: Marwan.Bukhari@mbht.nhs.uk) for reviewing the chapter.

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