Recent advances in the treatment of multiple sclerosis

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Abstract: Multiple sclerosis (MS) is a chronic inflammatory, immune-mediated, demyelinating, neurodegenerative disorder of the central nervous system. Despite the lack of an etiologic factor, it has been consistently demonstrated that the immune system plays a crucial role in the pathogenesis of MS. The traditional description of immunopathogenesis of MS suggests a preferential CD4+ TH1 cell activity causing tissue damage by the release of pro-inflammatory cytokines and subsequent demyelination and axonal loss. Recent evidence, however, suggests that other immune cells including TH17 cells, CD8+ effector T cells, CD4+ CD25+ regulatory T cells, and B cells may play a prominent role in MS immunopathology. A better understanding of the molecular and cellular components of the immunopathogenesis of MS is allowing the development of novel therapies.

Keywords: multiple sclerosis, immunopathology, oral medication, clinical trial

Introduction
Multiple sclerosis (MS) is a leading cause of disability among young adults in North America and Europe. Approximately 2.5 million people worldwide and 400,000 Americans have MS.1 Owing to the strong evidence suggesting immune-mediated mechanisms involved in the pathogenesis of the disease, MS therapies have been developed to modulate or suppress inflammatory responses. Current US Food and Drug Administration (FDA) approved treatments for MS include immunomodulators (interferon [IFN]-β, glatiramer acetate), or immunosuppressants (mitoxantrone, Tysabri®). These therapies are partially effective, with a wide range of tolerability and safety profiles. Considering the heterogeneity observed with respect to the clinical, radiographic, morphological, and genetic features of the disease, it is possible that various immunological mechanisms are involved in the pathogenesis of the disease. Besides the description of complex orchestrated immune-mediated mechanisms involved in the pathogenesis of the disease, recent pathological and magnetic resonance imaging (MRI) studies support the presence of diffuse tissue injury and neuronal degeneration extending beyond the classical white matter lesions.2,3 Therefore, it is imperative in the future to promote the development of individualized therapeutic strategies targeting both inflammatory and degenerative processes. Recent advances in the understanding of molecular and cellular components involved in the inflammatory and immune responses in MS have generated a new armamentarium of immune-based therapies, including several oral therapies in late-phase clinical trials and/or under review by the FDA for relapsing–remitting MS (RRMS). Here we review the most promising emerging therapies in development for the treatment of MS.
Elevated CD56Bright
Daclizumab

Immune sequestration
Fingolimod

Lymphocyte-targeted therapies
Cladribine
Teriflunomide

Cell proliferation
Cladribine
Teriflunomide

Antibody-dependent cell lysis
Alemtuzumab
Rituximab

Figure 1 Proposed mechanism of action of selected emerging therapies in MS.

Oral medications

Oral cladribine

Cladribine is a purine nucleoside analog engineered to exploit the specific enzymatic degradation of deoxynucleotides by adenosine deaminase (ADA) present in lymphocytes. Cladribine is phosphorylated intracellularly by deoxycytidine kinase to its active form, 2-chloro-2'-deoxyadenosine triphosphate. Because cladribine is resistant to ADA, it gets incorporated into DNA of lymphocytes and monocytes, and subsequently interrupts DNA replication, causing DNA damage and cell death. In addition, cladribine possesses epigenetic properties, by inhibiting S-adenosyl homocysteine hydrolase and DNA methylation. Initially, cladribine was thought to cause a selective and long-lasting reduction of activated CD4+ T lymphocytes with relatively mild and transient effects on monocytes and neutrophils. Recent data from a subset of patients participating in a Phase III randomized, placebo-controlled trial has demonstrated that cladribine produces a significant change in multiple peripheral blood subsets including an equal reduction in CD4+ and CD8+ T lymphocytes, a moderate reduction in CD56+ natural killer (NK) cells, and a rapid and severe reduction in B lymphocytes. These findings suggest that cladribine may act at multiple levels to modulate the immune system in relapsing disease: elimination of self-reactive helper and cytotoxic T cell clones, augmentation of NK regulation of autologous T lymphocytes, and reduction of effector B cells.

Previous studies evaluating parenteral cladribine in 257 MS patients showed a significant reduction in the number and volume of T1 gadolinium-enhancing lesions, accumulation of T2 lesion volume, relapse rate, and disability progression. A recent placebo-controlled Phase III trial using oral cladribine (3.5 or 5.25 mg/kg in the first year, and 3.5 mg/kg in the second year) in patients with RRMS showed a significant reduction in annualized relapse rates at 2 years compared with placebo (55%–58%), and reduction in the risk of progression to disability (hazard ratio for the 3.5-mg group, 0.67; 95% confidence interval [CI]:
initiated. Placebo-controlled and active comparator Phase III trials were recently completed.13,14 In the placebo-controlled Phase III trial, subjects using 0.5 mg of oral fingolimod demonstrated a significant reduction of annualized relapse rate, risk of accumulation of clinical disability, and MRI-related outcomes. In the active comparator study, oral fingolimod (0.5 and 1.25 mg daily) demonstrated superior efficacy versus intramuscular (IM) IFN (30 µg IM weekly) on relapse rate and MRI outcomes. Adverse reactions in both trials included bradycardia, macular edema, nasopharyngitis, dyspnea, headaches, diarrhea, and nausea. Three cases of basal cell carcinoma, three cases of squamous cell carcinoma, and one case of melanoma were reported. Two fatalities associated with varicella zoster and herpes virus infection and a single case of hemorrhagic encephalitis were also reported with an unclear causal relationship.

Laquinomod

Laquinomod (quinoline-3-carboxamid) is a once-daily, orally administered immunomodulatory compound that is being developed as a disease-modifying treatment for RRMS. In the experimental autoimmune encephalomyelitis animal model, laquinomod caused a dose-dependent reduction in disease severity, with decreased CD4+ T cell and macrophages infiltrates in spinal cord without resulting in significant systemic immunosuppression.15 Anti-inflammatory properties of laquinomod are thought to be secondary to a shift to a Th2/Th3 cytokine pattern. In addition it has been suggested that laquinomod may cause a downregulation of major histocompatibility complex class II gene transcription factors, stimulate neurotrophin (NT)-3, NT-4, and brain derived neurotrophic factor (BDNF) secretion, activate interleukin (IL)-4 signaling in CD4+ cells, promote apoptosis of CD8+ T and B cells, and suppress the metabolic activity of CD14+ and NK cells. A recent Phase IIb study of 306 patients demonstrated that an oral daily 0.6 mg dose of laquinomod significantly reduced MRI disease activity by a median of 60% versus placebo in RRMS patients. The majority of the patients that have participated in the trial are now receiving treatment with laquinomod in a continued open-label extension trial.16 An ongoing Phase III trial is currently evaluating efficacy of laquinomod (0.6 mg daily) in patients with RRMS.17

Laquinomod appears to be well tolerated, with only some transient and dose-dependent increases in liver enzymes. A case of Budd–Chiari syndrome occurred after one month of exposure in a patient with underlying hypercoagulability. Unlike its precursor substance, linomide, no cases of serositis...
or myocardial infarction have been reported so far in patients receiving laquinimod.

**BG00012 (oral fumarate)**

BG00012 is an oral formulation of dimethyl fumarate. The exact mechanism of action is not known but is thought to inhibit immune cells by stimulating the expression of anti-inflammatory cytokines such as IL-10, IL-4, and IL-5, resulting in a shift from a Th-1 (pro-inflammatory) to a Th-2 (anti-inflammatory) T cell response. In addition, BG00012 may provide a neuroprotective therapeutic effect by inducing phase II detoxification genes, and upregulation of the phase II detoxification enzyme NAD(P)H:quinone oxidoreductase-1 (NQO-1). A multicenter, randomized, placebo-controlled clinical trial of oral BG00012 involving 257 people with RRMS, receiving various doses of BG00012, demonstrated a dose-dependent reduction in active inflammation on MRI scans. Tolerability was good overall, with adverse events more commonly occurring in the treatment groups, including abdominal pain and flushing. Two Phase III trials are currently ongoing, evaluating whether BG00012 is effective in reducing the proportion of relapses, decreasing the number of brain lesions, and slowing time to progression. Estimated completion dates are in December 2010.

SB-683699 is an oral medication that is thought to inhibit leukocyte trafficking through the BBB by antagonism of alpha 4 integrins. A Phase II trial in patients with RRMS evaluating safety and efficacy of SB-683699 on the development of new brain lesion formation at 6 months as detected by MRI scan as well as accumulation of disability will provide results soon.

**Teriflunomide**

Teriflunomide is a dihydroorotate dehydrogenase inhibitor, a mitochondrial enzyme crucially involved in pyrimidine synthesis. Because activated lymphocytes depend on de novo pyrimidine synthesis, pyrimidine depletion results in inhibition of immune-cell proliferation. There is also some evidence from in vitro studies suggesting that teriflunomide may induce Th-2-mediated anti-inflammatory cytokine activation.

Oral teriflunomide was tested in a randomized, double-blind, placebo-controlled Phase II study. Patients with relapsing forms of MS were randomized to receive placebo, teriflunomide 7 mg, or teriflunomide 14 mg per day for 36 weeks. Teriflunomide demonstrated a dose-dependent reduction in the number of T₁-enhancing lesions. Teriflunomide was generally safe and well tolerated. Adverse effects included nasopharyngitis, alopecia, nausea, limb pain, diarrhea, and arthralgia. Hepatic necrosis and pancytopenia have been reported in patients with rheumatoid arthritis who were taking teriflunomide. A 2-year, double-blind, placebo-controlled Phase III study in relapsing MS is in progress. The primary outcome measure is relapse rate. Other ongoing or planned studies of teriflunomide include a Phase II study of combination with IFN-β, a Phase II study of combination with glatiramer acetate, and a placebo-controlled Phase III trial in clinically isolated syndrome.

**Parenteral medications**

**Alemtuzumab**

Alemtuzumab is a humanized monoclonal antibody directed against CD52, a cell surface receptor expressed on lymphocytes, NK cells, monocytes, and macrophages. Binding of the monoclonal antibody to B and T lymphocytes results in antibody-dependent cell lysis and a prolonged elimination (up to 16 months) of cells from the bone marrow and blood. Interestingly, the capacity of immune cells to regenerate remains intact after repetitive cycles of alemtuzumab, but immature forms of B cells (transitional type I cells), possibly driven by high BAFFs levels, are seen early in the course of immune reconstitution. This event could explain the frequent development of antibody-mediated autoimmune disorders with alemtuzumab therapy. Alemtuzumab is currently US Food and Drug Administration (FDA) approved to treat B-cell chronic lymphocytic leukemia.

Studies of alemtuzumab in the treatment of patients with RRMS and secondary progressive MS (SPMS) have shown efficacy in the suppression of annualized relapse rate but with variable results in preventing progression of disability, depending on stages of the disease. In a recent Phase II randomized blinded clinical trial (IV at 12 mg per day or 24 mg per day for 5 consecutive days during the first month and on 3 consecutive days at 12 and 24 months), alemtuzumab significantly reduced the rate of sustained accumulation of disability by 71% with a 74% reduction on the annualized rate of relapse when compared with IFN-β-1a in patients with early RRMS. Over 80% of patients receiving alemtuzumab remained relapse free at 36 months. Reduction in T₂ lesion volume and total brain volume analysis was greater in the alemtuzumab group. Adverse events in the alemtuzumab group included autoimmunity (thyroid disorders and immune thrombocytopenic purpura) and infections. About one third of patients develop antibodies against the thyrotropin receptor and subsequent autoimmune hyperthyroidism.
Currently there are two Phase III trials comparing two annual cycles of intravenous alemtuzumab to three times weekly IFN-β-1a in treatment naïve patients with RRMS.29,30 These studies are expected to be completed in 2011–2012.

Rituximab is a chimeric antibody directed against the human CD20 phosphoprotein that is present on all B cell lineages except for stem cells, pro-B cells, and plasma cells. Rituximab acts by binding to B cell lymphocytes, initiating complement-mediated B cell lysis. The standard dose of rituximab is 375 mg/m² given weekly for 4 weeks, or a fixed dose of 2 g divided in two infusions of 1 g each, 2 weeks apart. Following an intravenous administration of 2 g rituximab, CD20+ B cells are rapidly depleted and remain undetectable for up to 6 months longer.

Rituximab has shown efficacy in the treatment of patients with RRMS. A recently completed Phase II randomized clinical trial using a biweekly 1000 mg dose of rituximab in RRMS patients demonstrated a 91% reduction on the number of gadolinium-enhancing lesions on MRI scans, as well as significant reduction in clinical relapses. Mild infusion-related complications were seen in most patients, but serious adverse reactions were rare.14 In a recent trial of patients with primary progressive MS, rituximab appeared to have efficacy only in young patients with signs of active inflammation on MRI scans.33 Five cases of progressive multifocal leukoencephalopathy have been recently reported in patients receiving rituximab for the treatment of rheumatoid arthritis and systemic lupus erythematosus. However, those patients were receiving additional immunosuppressants. Ocrelizumab, a humanized monoclonal antibody against human CD20, is currently under investigation in a Phase II trial evaluating efficacy and safety in patients with RRMS. Estimated completion of the trial is 2012.32

Daclizumab is a humanized mouse monoclonal antibody that binds to the alpha subunit of the IL-2 receptor. This receptor is present on activated T and B cells and is crucial for T cell proliferation and activation. The clinical benefit of daclizumab has been linked to significant expansion of immunoregulatory CD56 NK cells, and subsequent downregulation of adaptive T cell responses (CD4+ and CD8+ T cells).33 In a first open-label study with intravenous daclizumab (1 mg/kg dose), five patients with SPMS and six patients with RRMS demonstrated a decrease of contrast-enhancing lesions by 78% and in a relapse rate by 81% compared with baseline.34 A recent open-label Phase II trial using subcutaneous daclizumab (2 mg/kg doses) in MS patients with inadequate response to IFN therapy demonstrated a 72% reduction in the number of new or enlarged contrast-enhancing MRI lesions at week 24 compared with patients receiving IFN-β alone. Because type 1 IFNs are known to also enhance NK-cell function, the question of possible synergism between IFN and daclizumab therapy is possible.35 Skin rash, chest discomfort, headaches, lymphopenia, generalized lymphadenopathy, and transient elevation of liver function tests and bilirubin level were reported. Daclizumab is already in clinical use to prevent rejection of kidney transplants. A multicenter Phase II trial investigating a subcutaneous formulation of daclizumab (DacHyp) monotherapy is ongoing.

CTLA4Ig is a chimeric fusion protein that prevents T cell activation by binding the B7–1 and B7–2 receptors. 33 In a first open-label study with intravenous daclizumab, 72% of patients tested were found to have evidence of a significant increase in the number of immunoregulatory CD56 NK cells, and subsequent downregulation of adaptive T cell responses (CD4+ and CD8+ T cells). In a recent open-label trial evaluating efficacy and safety in patients with RRMS, estimated completion of the trial is 2011–2012.

### Table 1: Summary of characteristics, efficacy, and safety information on selected MS therapies

<table>
<thead>
<tr>
<th>Mechanism of action</th>
<th>Route/dose</th>
<th>Results</th>
<th>AEDs</th>
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<tbody>
<tr>
<td>Alemtuzumab</td>
<td>Anti-CD52 IV 12 mg dose/y</td>
<td>75% ↓ SAD (12 mg dose), 74% ↓ RR</td>
<td>Skin rash, chest discomfort, headaches, lymphopenia</td>
</tr>
<tr>
<td>Daclizumab</td>
<td>Anti-CD25/anti-IL2, NK CD56 SQ 2 mg/kg (Q2w)</td>
<td>72% ↓ CEL in RRMS patients</td>
<td>Infusion reaction, allergies</td>
</tr>
<tr>
<td>Rituximab</td>
<td>Anti-CD20 IV (1 g dose, Q2w, ×2)</td>
<td>91% ↓ CEL in RRMS</td>
<td>Bradycardia, nasopharyngitis, dyspnea, headaches, diarrhea and nausea, hemorrhagic encephalitis, skin cancer</td>
</tr>
<tr>
<td>Rituximab</td>
<td>Anti-CD20 IV (1 g dose, Q2w, ×2)</td>
<td>91% ↓ CEL</td>
<td>Bradycardia, nasopharyngitis, dyspnea, headaches, diarrhea and nausea, hemorrhagic encephalitis, skin cancer</td>
</tr>
<tr>
<td>Fingolimod</td>
<td>Sphingosine-1-phosphate (S1P) analog Oral (0.5 and 1.25 mg)</td>
<td>54%–60% ↓ ARR</td>
<td>Skin rash, chest discomfort, headaches, lymphopenia</td>
</tr>
<tr>
<td>Laquinimod</td>
<td>Anti-inflammatory Oral 0.6 mg/d</td>
<td>40% ↓ CEL</td>
<td>Skin rash, chest discomfort, headaches, lymphopenia</td>
</tr>
<tr>
<td>Fumaric acid (BG12)</td>
<td>Anti-inflammatory/ neuroprotective Oral (120–240 mg TID)</td>
<td>69% ↓ CEL</td>
<td>Infusion reaction, allergies</td>
</tr>
<tr>
<td>Teriflunomide</td>
<td>Inhibition of immune cell proliferation Oral (7 and 14 mg dose)</td>
<td>61% ↓ CEL</td>
<td>Skin rash, chest discomfort, headaches, lymphopenia</td>
</tr>
<tr>
<td>Cladribine</td>
<td>Purine nucleoside analog Oral (3.5 and 5.25 mg/kg total dose)</td>
<td>58% ↓ RR, 43% disease free</td>
<td>Skin rash, chest discomfort, headaches, lymphopenia</td>
</tr>
</tbody>
</table>

**Abbreviations:** ARR, annualized relapse rate; CEL, contrast enhancing MRI lesions; HA, headaches; HZV, herpes zoster virus; ITP, immune thrombocytopenic purpura; IV, intravenous; LFT, liver function test; RR, relapse rate; RRMS, relapsing remitting multiple sclerosis; SAD, sustained accumulation of disability; SQ, subcutaneous.
A recent Phase I clinical trial showed that intravenous administration of CTLA4Ig was well tolerated in patients with MS, and most adverse events were rated as mild. Lymphadenopathy, urinary tract infections, headaches, blurred vision, and upper respiratory tract infections were most frequently reported. Immunologic assessment of the patients showed a reduction in myelin basic protein (MBP) proliferation within 2 months of infusion and decreased IFN-γ production by MBP-specific lines. No significant changes in clinical or MRI parameters were observed during the duration of the study.36

**Comments**

The etiology of MS remains unclear; however, there has been a significant progress in our knowledge of the basic pathophysiological mechanisms involved in the disease process. Clinicoepidemiological studies suggest a high level of heterogeneity among MS patients. In addition, recent studies suggest that endogenous expression of certain cytokines might impact on the clinical efficacy of currently available therapies.37 Considering the complex mechanisms of action of current and emerging therapies, it is imperative to promote the development of biomarkers that might have predictive value in identifying future response to various therapies.

**Disclosure**

The authors report no conflicts of interest in this work.

**References**

17. Teva Pharmaceutical Industries. A multinational, multicenter, open-label, single-assignment extension of the MS-LAQ-301 Study, to evaluate the long-term safety, tolerability and effect on disease course of daily oral laquinimod 0.6 mg in subjects with relapsing multiple sclerosis. NCT00988052.


