

# Benefit–Risk Profile of Sphingosine-1-Phosphate Receptor Modulators in Relapsing and Secondary Progressive Multiple Sclerosis

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**Abstract** Since the approval of fingolimod, several selective sphingosine-1-phosphate receptor modulators have entered clinical development for multiple sclerosis. However, side effects can occur with sphingosine-1-phosphate receptor modulators. By considering short-term data across the drug class and longer term fingolimod data, we aim to highlight the potential of sphingosine-1-phosphate receptor modulators in multiple sclerosis, while offering reassurance that their benefit–risk profiles are suitable for long-term therapy. Short-term fingolimod studies demonstrated the efficacy of this drug class, showed that cardiac events upon first-dose administration are transient and manageable, and showed that serious adverse events are rare. Early-phase studies of selective sphingosine-1-phosphate receptor modulators also show efficacy with a similar or improved safety profile, and treatment initiation effects were reduced with dose titration. Longer term fingolimod

studies demonstrated sustained efficacy and raised no new safety concerns, with no increases in macular edema, infection, or malignancy rates. Switch studies identified no safety concerns and greater patient satisfaction and persistence with fingolimod when switching from injectable therapies with no washout period. Better outcomes were seen with short than with long washouts when switching from natalizumab. The specific immunomodulatory effects of sphingosine-1-phosphate receptor modulators are consistent with the low observed rates of long-term, drug-related adverse effects with fingolimod. Short-term data for selective sphingosine-1-phosphate receptor modulators support their potential effectiveness in multiple sclerosis, and improved side-effect profiles may widen patient access to this drug class. The long-term safety, tolerability, and persistence profiles of fingolimod should reassure clinicians that sphingosine-1-phosphate receptor modulators are likely to be suitable for the long-term treatment of multiple sclerosis.

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## Key Points

Sphingosine-1-phosphate receptor modulators are a new class of oral disease-modifying therapy for multiple sclerosis (MS); this review summarizes both short-term efficacy and safety data available for this drug class, and long-term data for fingolimod.

Short-term data demonstrate the clinical effectiveness on relapses, magnetic resonance imaging lesions, and disability progression of this drug class in MS, while serious adverse events are rare, and side effects at first dose are transient and manageable.

Fingolimod is an effective MS therapy in the long term, and neither its long-term use nor switching to fingolimod from other treatments increases the risk of adverse events.

The positive benefit–risk profile of sphingosine-1-phosphate receptor modulators should reassure clinicians that this drug class is suitable for the long-term treatment of MS.

## 1 Introduction

Several selective sphingosine-1-phosphate (S1P) receptor modulators are in clinical development for the treatment of patients with relapsing forms of multiple sclerosis (RMS) or secondary progressive multiple sclerosis (SPMS) [1–3]. These compounds were developed following the success of fingolimod [1–5], a functional antagonist of S1P receptor subtypes S1P<sub>1,3–5</sub>, and the first oral disease-modifying therapy (DMT) approved for the treatment of RMS in USA and of relapsing-remitting multiple sclerosis in Europe [6–8]. Sphingosine-1-phosphate receptor modulators under development include ponesimod, siponimod, ozanimod, ceralifimod, GSK2018682, and amiselimod [9–11].

Sphingosine-1-phosphate receptor modulators have not been shown to be effective in primary progressive multiple sclerosis. A phase III study of fingolimod in patients with primary progressive multiple sclerosis showed a small but non-significant improvement in confirmed disability progression (CDP) with fingolimod over placebo and similar percentage brain volume loss (BVL) in the two treatment groups [12]. This review therefore focuses on the role of S1P receptor modulators in RMS and SPMS.

Other articles have reviewed the mechanisms of action of S1P receptor modulators [2, 11, 13, 14]; thus, here we provide an overview before summarizing short-term

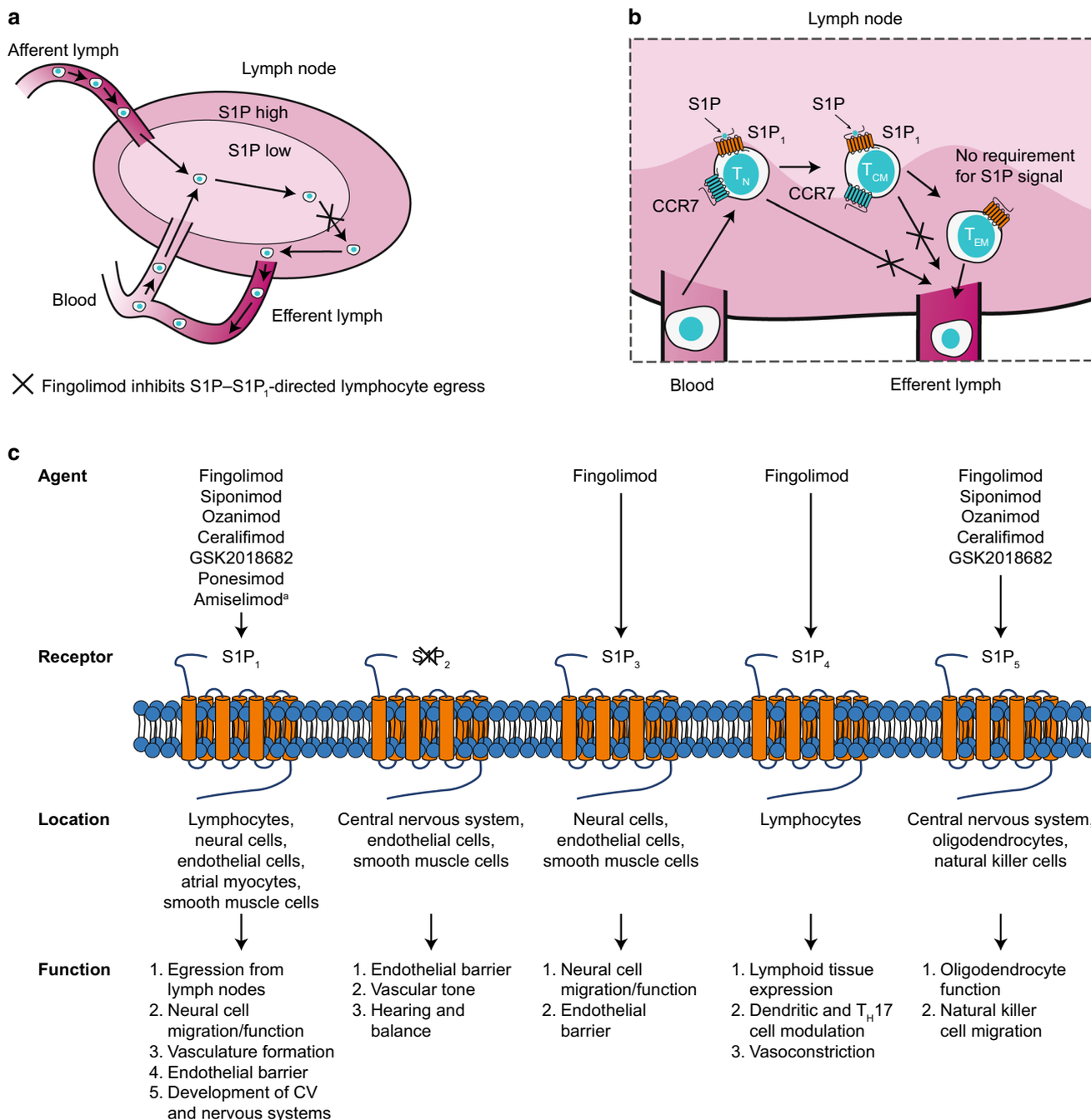
efficacy and safety data ( $\leq 2$  years) across the drug class, and longer term data ( $> 2$  years) available from studies of fingolimod. Real-world and observational studies as well as phase II and III clinical trials are considered to understand outcomes among patients potentially under-represented in controlled trials. Here, we aim to illustrate the potential of S1P receptor modulators in the treatment of patients with RMS and SPMS, and provide reassurance that their benefit–risk profiles are suitable for long-term therapy. Studies are summarized in Online Resource 1 of the Electronic Supplementary Material (ESM) [e1–e40].

## 2 Mechanism of Action of Sphingosine-1-Phosphate (S1P) Receptor Modulators

Sphingosine-1-phosphate is a bioactive phospholipid regulating a range of cellular processes, including immunity, inflammation, angiogenesis, heart rate, smooth muscle tone, cell differentiation, cell migration and survival, calcium homeostasis, and endothelium integrity. Its effects are mediated by the G-protein-coupled receptor subtypes S1P<sub>1–5</sub> [3, 11]. The pattern of expression and roles of different S1P receptor subtypes are summarized in Fig. 1 [11, 15]. S1P<sub>1</sub>, S1P<sub>2</sub>, and S1P<sub>3</sub> are ubiquitous [11] but most abundant in immune, cardiovascular, and central nervous system cells [3]; S1P<sub>4</sub> is mostly expressed in lymphoid tissues and the lungs [3, 11]; and S1P<sub>5</sub> in the central nervous system (on oligodendrocytes), skin, and spleen [3, 11]. S1P<sub>1</sub> signaling mediates the trafficking of lymphocytes, particularly their egress from lymph nodes and migration into the blood and target tissues, apparently without inhibiting their activation, proliferation, or effector functions [3, 8, 11, 15–17].

### 2.1 Fingolimod

Fingolimod is a structural analog of sphingosine [8] and, after phosphorylation, is a functional antagonist of S1P<sub>1,3–5</sub> [11], which acts reversibly to retain circulating central memory T cells and naïve T cells in the lymph nodes, thereby reducing the migration of autoreactive lymphocytes into the central nervous system [8]. Effects of fingolimod on B cells and natural killer cells have also been reported [8], and some experimental evidence exists for a direct neuroprotective effect of fingolimod on brain cells (reviewed in Hunter et al. [8] and Jeffery et al. [15]). The therapeutic effects of fingolimod are primarily mediated via S1P<sub>1</sub> on lymphocytes; however, certain adverse events (AEs) involving heart rate and atrioventricular (AV) conduction are also caused by the initial agonistic interaction with S1P<sub>1</sub>, while those involving macular edema or pulmonary function may be associated with its nonselective binding to other S1P receptors, such as S1P<sub>3</sub> [11, 15].



**Fig. 1** Mechanism of action of sphingosine-1-phosphate (S1P) receptor modulators. **a** The binding of S1P receptor modulators to S1P<sub>1</sub> on central memory T-cells (T<sub>CM</sub>) causes these cells to engulf their own S1P<sub>1</sub>, resulting in T<sub>CM</sub> that are unresponsive to S1P signals. Any new S1P receptors being produced inside the cell remain in a state of arrest until S1P receptor modulation is removed. Therefore, T<sub>CM</sub> do not leave the lymph node in response to S1P signals, and, by inhibiting the movement of T<sub>CM</sub> into the circulation, S1P receptor modulators prevent these autoreactive cells from migrating into the central nervous system. In contrast, the levels of peripheral effector memory T-cells (T<sub>EM</sub>) are largely unaffected by S1P receptor

modulators, thus preserving immunosurveillance and the capacity to respond to and contain locally invading pathogens. **b** Interaction of S1P receptor modulators with S1P receptor subtypes. **c** Interaction of fingolimod and selective S1P receptor drugs with S1P receptor subtypes. <sup>a</sup>Amiselimod is selective, but its selectivity is unknown. CCR7 C-C chemokine receptor type 7, CV cardiovascular, S1P<sub>1-5</sub> sphingosine-1-phosphate receptor subtypes 1–5, T<sub>H</sub>17 T helper cell 17, T<sub>N</sub> naïve T-cell. Parts **a** and **b** reproduced with permission from Jeffery et al. Expert Rev Neurother. 2016;16:31–44 [15]. Part **c** reproduced/adapted with permission from Subei and Cohen, CNS Drugs. 2015;29:565–75 [11]

## 2.2 Other S1P Receptor Modulators

Discovery of the receptor-binding characteristics of fingolimod has triggered a search for S1P receptor modulators with a greater selectivity for S1P<sub>1</sub>. Agents currently under development include ponesimod, siponimod, amiselimod, and ozanimod [11]; GSK2018682 has completed phase I trials [18], although further development seems unlikely [11], and a phase III study of ceralifimod was terminated prematurely, thus its future development is also uncertain [19, 20]. Ponesimod is an orally active, reversible, and selective S1P<sub>1</sub> modulator [21]; siponimod, ozanimod, and ceralifimod are orally active selective modulators of S1P<sub>1</sub> and S1P<sub>5</sub> [22–26]; GSK2018682 is a selective S1P<sub>1</sub> modulator [18] that has some activity at S1P<sub>5</sub>; and amiselimod is known to be a selective S1P receptor modulator, but the subtype selectivity is unknown [11]. Like fingolimod, the therapeutic effects of these compounds are primarily mediated by rapid internalization, degradation, and functional antagonism of S1P<sub>1</sub>, leading to lymphocyte sequestration in the lymph nodes; the lack of activity on S1P<sub>3–4</sub> may alleviate some side effects, including those at first dose. All of these compounds (apart from amiselimod) also have a shorter half-life and show a reduced time to lymphocyte recovery after treatment discontinuation compared with fingolimod, providing an improved safety profile for patients who need to interrupt medication for any reason. The downstream effects of the interactions between these compounds and different S1P receptor subtypes are shown in Fig. 1 [11, 15], and a summary of the S1P receptor modulator pharmacokinetics and pharmacodynamics is provided in Table 1 [6, 7, 18, 21, 22, 27–30].

## 3 Short-Term Efficacy (≤2 Years)

Randomized controlled studies [31–34] of fingolimod typically range from 6 months to 2 years in duration, whereas data published for ponesimod, siponimod, amiselimod, and ozanimod are from dose-ranging clinical studies of no more than 6 months' duration [21, 28, 35, 36]; short-term efficacy data from these studies are summarized in Online Resource 2 of the ESM [e1, e2, e4, e7, e20, e25, e28, e32].

### 3.1 Fingolimod

As of November 2016, approximately 184,000 patients with multiple sclerosis (MS) have been treated with fingolimod, with total patient exposure exceeding 397,000 patient-years [37]. In phase II and III studies [31–34], significant reductions in annualized relapse rate (ARR) and magnetic resonance imaging (MRI) markers of disease activity (gadolinium-enhancing [Gd+] and T2 lesion counts) were consistently observed with fingolimod compared with placebo or intramuscular (IM) interferon (IFN) β-1a. Significantly less BVL was observed with fingolimod than with placebo or IFNβ-1a in studies with a duration longer than 6 months, and significant reductions in the volume of T1-hypointense 'black holes' were seen with fingolimod compared with placebo. In the placebo-controlled studies, there were trends towards proportionally more patients remaining free from 3- and 6-month CDP with fingolimod than with placebo; this difference was statistically significant in the FREEDOMS trial [32].

**Table 1** Summary of sphingosine-1-phosphate receptor modulator pharmacokinetics and pharmacodynamics

	Pro-drug (requires phosphorylation in vivo)	$T_{\max}$ (h)	Time to lymphocyte count reduction (h)	Lymphocyte decrease from baseline (%)	$t_{1/2}$ (h)	Time to lymphocyte count recovery after treatment discontinuation (days)
Fingolimod [6, 7]	Yes	12–16	4–6	70	144–216	30–60
Ponesimod [21, 27]	No	2.5–4	6	50–70	32	7
Siponimod [22]	No	3–4.5	4–6	33–76	30	1–5
Ozanimod [28, 29]	No	6–8	6–12	34–68	19	2–3
GSK2018682 [18]	No	4–9	6–16	28–76	48–63	1–6
Amiselimod [30]	Yes	12–16	No data	60–66	380–420	49

$T_{\max}$  time to maximum plasma concentration,  $t_{1/2}$  elimination half-life

### 3.2 Other S1P Receptor Modulators

Trials of ponesimod, amiselimod, siponimod, and ozanimod have shown treatment effects on MRI disease activity and ARR in patients with RMS [21, 28, 35, 36]; however, CDP was not assessed in these studies. In patients receiving ponesimod, significant reductions in the numbers of Gd + lesions and combined unique active lesions were reported at all doses compared with placebo. Lower ARR (significant at 40 mg) and smaller cumulative numbers of T2 lesions (significant at 20 mg) were also reported with ponesimod than with placebo [21]. A dose-dependent numerical reduction in BVL was observed with ponesimod, although *p* values were not reported.

Amiselimod provided significant dose-dependent reductions in the mean total numbers of T1 Gd + lesions compared with placebo from weeks 8 to 24 in patients with RMS [36]. The mean total number of new or enlarged T2 lesions from weeks 8 to 24 was also significantly reduced with amiselimod 0.2 and 0.4 mg compared with placebo. Annualized relapse rates were lower with amiselimod 0.2 and 0.4 mg than with placebo, although the difference was significant only in the 0.4-mg group. Brain volume loss after 24 weeks of treatment was similar in the amiselimod and placebo groups, although reductions in gray matter volume were significantly smaller with all amiselimod doses than with placebo.

After 3 months of siponimod treatment, lesion activity was reduced compared with placebo; significant relative reductions were observed in the numbers of combined unique active lesions (doses  $\geq 0.5$  mg), new Gd + lesions (doses  $\geq 0.5$  mg), and T2 lesions (doses  $\geq 1.25$  mg). At 6 months, there were significant relative reductions in the number of new Gd+ and T2 lesions with siponimod compared with placebo at doses of  $\geq 0.5$  and  $\geq 2$  mg, respectively. Over 6 months, ARRs were generally lower with siponimod than with placebo, but differences were significant only in the 2-mg group [35]. In the extension study, reductions in Gd + lesions and ARR were sustained over 2 years with  $\geq 1.25$  mg; at the 2-mg dose, 57.7% of patients remained free from combined unique active lesions at 2 years [10]. Positive outcomes with ozanimod have also been reported, with significantly lower cumulative numbers of Gd+ and T2 lesions during weeks 12–24 at both the 0.5- and 1-mg doses compared with placebo. Over 24 weeks, numerically lower ARRs were also reported at both doses compared with placebo [28].

Siponimod has been evaluated in patients with SPMS. In the phase III trial of siponimod 2 mg, the risk of 3- and 6-month CDP was significantly reduced by 21 and 26%,

respectively, compared with placebo [38]. Full results from this study have not been published to date.

## 4 Short-Term Safety ( $\leq 2$ Years)

Although the therapeutic effects of S1P receptor modulators are primarily mediated via binding to S1P<sub>1</sub> on lymphocytes, interactions with this and other S1P receptor subtypes may lead to AEs, for example decreases in heart rate and AV conduction delays after the first dose(s), mild increases in blood pressure, macular edema, and bronchoconstriction. Short-term safety data are summarized in Online Resource 3 of the ESM [e1, e2, e4, e7, e16, e17, e20, e21, e24, e25, e28, e32].

### 4.1 Fingolimod

In the phase III FREEDOMS, FREEDOMS II, and TRANSFORMS trials [32–34], AEs occurred in similar proportions of patients receiving fingolimod 0.5 mg (86–98%), IFN $\beta$ -1a (91.6%), and placebo (92–97%); similarly, serious AEs (SAEs) and discontinuations owing to AEs occurred in 5.8–15% and 3.7–18% of patients, respectively, and proportions were similar across treatment groups in each study [32–34]. First-dose effects of fingolimod treatment were consistently mild, with transient decreases in heart rate of 8.0–8.5 bpm that started to attenuate after 6 h. Bradycardia was reported in 0.7–2.1% of patients receiving fingolimod 0.5 mg; all cases resolved within 24 h. Second-degree AV block was rarely reported and was absent in one study [33]; where reported, it occurred in 0.2–0.7% of patients. Elevated alanine aminotransferase levels were found in 8.0–8.5% of patients, but all were asymptomatic. Overall incidences of infection were similar in patients receiving fingolimod 0.5 mg, placebo, or IFN $\beta$ -1a (42–74%); however, herpes virus, upper respiratory tract, and urinary tract infections were consistently more common with fingolimod than with controls. Macular edema was reported in 0.5–1.0% of patients receiving fingolimod 0.5 mg, but all affected patients were asymptomatic or had stable visual acuity; only one case of macular edema was classified as an SAE. Malignancies were reported at low frequencies across the phase III studies, with basal cell carcinoma being most common, affecting 0.7–3.0% of patients receiving fingolimod 0.5 mg. Incidences of AEs, SAEs, and discontinuations owing to AEs in real-world studies of fingolimod were generally lower than, or at the lower range of, those observed in the phase III trials [39–43]. In a phase II study, overall safety profiles of fingolimod 1.25 and 5.0 mg were similar to that seen with fingolimod 0.5 mg [31].

## 4.2 Other S1P Receptor Modulators

Safety data from short-term studies of ponesimod, siponimod, amiselimod, and ozanimod are broadly similar to those from fingolimod studies [10, 21, 28, 35, 36]; AEs occurred at similar rates for active treatment and placebo in each of the trials, the most common AEs, such as headache, nasopharyngitis, dizziness, and influenza, occurring at rates similar to those reported with fingolimod. Serious AEs in patients receiving ponesimod and siponimod were somewhat more frequent than with placebo, as were AEs causing discontinuation (ponesimod, 5.3–13.4%, placebo, 2.5%; siponimod, 12–20% placebo, 4.0%) [21, 35]. With amiselimod, incidences of AEs causing discontinuation and of SAEs were similar in the amiselimod and placebo groups [36]. With ponesimod and siponimod, first-dose effects (decreased heart rate, bradycardia, and second-degree AV block) and elevated alanine aminotransferase levels occurred at rates similar to those seen with fingolimod; however, cardiac events were reported as reasons for discontinuation in 2.6% of patients receiving ponesimod and 3.4% of those receiving siponimod [21, 35], and second-degree AV block was the only SAE seen with siponimod that was reported in at least two patients in the same treatment group [21, 35].

In the siponimod extension study, dose titration effectively reduced the first-dose effects on heart rate seen with higher doses, with no cardiac SAEs reported [10]. Proportionally more cases of macular edema were classified as SAEs with ponesimod than with siponimod, and these cases resulted in treatment discontinuation. Incidences of infections were similar among patients treated with placebo, ponesimod, or siponimod, with urinary and upper respiratory tract infections being most common; unlike with fingolimod treatment, no cases of herpes virus infection were reported following treatment with ponesimod or siponimod [21, 35]. Malignancies occurred at low rates with ponesimod and siponimod [10, 21, 35].

The safety profiles of amiselimod and ozanimod differed somewhat from those reported for the other compounds. First-dose effects were not observed with amiselimod [36]. Mean heart rate remained stable during the 6 h after initial dosing and throughout the study, and there was no clinically relevant bradycardia, sinus pause, or AV block in any amiselimod group. One episode each of second-degree AV block and non-sustained ventricular tachycardia were reported as SAEs with amiselimod; however, both patients were asymptomatic, and these events were not considered clinically significant by the cardiologist on the safety monitoring board [36]. Moreover, no cases of macula edema were reported with amiselimod. Incidences of infections were similar among patients treated with placebo and amiselimod (26–32%), with nasopharyngitis and upper respiratory tract

infections being most common. Headache was reported more frequently in each of the amiselimod groups (10%) than with placebo (4%); frequencies of cardiac disorders were low and similar across treatment groups (4–8%) [36]. Liver enzyme abnormalities led to discontinuations in three patients (one each in the placebo [1%], amiselimod 0.2-mg [1%], and amiselimod 0.4-mg [1%] groups), and slight dose-dependent increases in mean values for alanine aminotransferase, aspartate aminotransferase, and  $\gamma$ -glutamyltransferase were seen with amiselimod; however, no symptoms were associated with these abnormal values, thus they were not considered to be clinically significant [36]. With ozanimod, there were no discontinuations owing to an AE, and no cases of macular edema, notable infections, notable cardiovascular AEs, or malignancies; and with dose titration, heart rate was reduced at first dose by  $\leq 2$  bpm [28].

## 5 Long-Term Efficacy (>2 Years)

The efficacy benefits of fingolimod reported during the 2-year FREEDOMS trial were sustained in the 2-year FREEDOMS extension study [44]. In the continuous-fingolimod patient groups, the ARR was lower ( $p < 0.0001$ ), BVL was reduced ( $p < 0.05$ ), and a greater proportion of patients were free from 3-month CDP ( $p < 0.05$ ) compared with the placebo–fingolimod switch-patient groups [44]. In the placebo–fingolimod switch groups, ARR was lower ( $p < 0.001$  for both groups), and BVL was reduced ( $p < 0.01$  for the placebo–fingolimod 0.5 mg group) after switching to fingolimod [44]. Similarly, the low levels of clinical and MRI markers of disease activity reported with fingolimod in the FREEDOMS II trial were maintained in the extension among patients who continued on treatment [45]. From baseline to the end of the study, ARR was significantly lower in the continuous-fingolimod groups than in the placebo–fingolimod switch groups ( $p < 0.001$ ); ARR, MRI disease activity, and BVL rates remained low with continuous fingolimod treatment in the trial extension [45].

In the TRANSFORMS extension study, effects of fingolimod were maintained in the long term (for up to 4.5 years), with low rates of disease activity and improvements in efficacy observed after patients switched from IFN $\beta$ -1a IM to fingolimod [46]. From baseline to the end of study, ARR was significantly lower in patients receiving continuous fingolimod than in patients in the IFN $\beta$ -1a–fingolimod switch group (0.17 vs. 0.27, respectively; 35% reduction in relapse risk;  $p < 0.001$ ) [46]. In a post hoc analysis, the proportion of patients with no evidence of disease activity (relapses, 3-month CDP, MRI activity; no evidence of disease activity-3) increased by approximately 50% in the 1st year after switching to fingolimod (from 44.3 to 66.0%) [46].

After 5 years in the ongoing LONGTERMS study, a single-arm open-label extension to various phase II, III, and IIIb trials of fingolimod (Fig. 2) [47], clinical disease activity remained low in patients treated with fingolimod [48]. Annualized relapse rate was low during the 1st year and remained low during each subsequent year up to interim analysis [48]. Mean Expanded Disability Status Scale scores remained stable, with most patients not reaching an Expanded Disability Status Scale score higher than 3 after receiving fingolimod [48]. Overall, most patients remained free from 6-month CDP in the FREEDOMS (78%) and TRANSFORMS (75%) cohorts [48].

Results of the 36-month interim analysis [49] of the PANGAEA study, a 5-year prospective study designed to collect real-world efficacy, safety, and pharmaco-economic data from patients receiving fingolimod in Germany [50], supported the positive efficacy profile demonstrated by fingolimod in the phase III clinical trials and extensions [49]. A mean baseline ARR of 1.5 was reduced to 0.43, 0.41, and 0.37 at 12, 24, and 36 months, respectively. In the first, second, and third years of the study, 67.9, 68.3, and 70.1% of patients, respectively, were relapse-free [49]. The mean Expanded Disability Status Scale score (3.0) was stable over 36 months, and 70.5% of patients had no clinical disease activity in year 3 [49]. Treatment satisfaction improved by 3 points over 24 months [49].

## 6 Long-Term Safety Data (>2 Years)

In the extension [44] to the 2-year FREEDOMS trial [32], long-term fingolimod treatment was well tolerated, and no new safety indications were observed [44]. Adverse events associated with treatment initiation were seen in the switch group, but the overall incidences and patterns of AEs (91.7–95.5%) and SAEs (7.1–11.7%) were similar across groups [44]. Cardiac effects associated with initiation of fingolimod were transient; blood pressure remained stable in the continuous-treatment groups [44].

In the 2-year extension to FREEDOMS II [45, 51], the safety profile of fingolimod was consistent with that observed in the core study. The incidences of AEs were similar across the continuous-fingolimod and treatment-switch groups (85.3% and 87.9%) [51]; macular edema was confirmed in one patient in each switch group [51], and there was one reported occurrence of squamous cell carcinoma and one of basal cell carcinoma in the continuous-fingolimod 0.5-mg group [51].

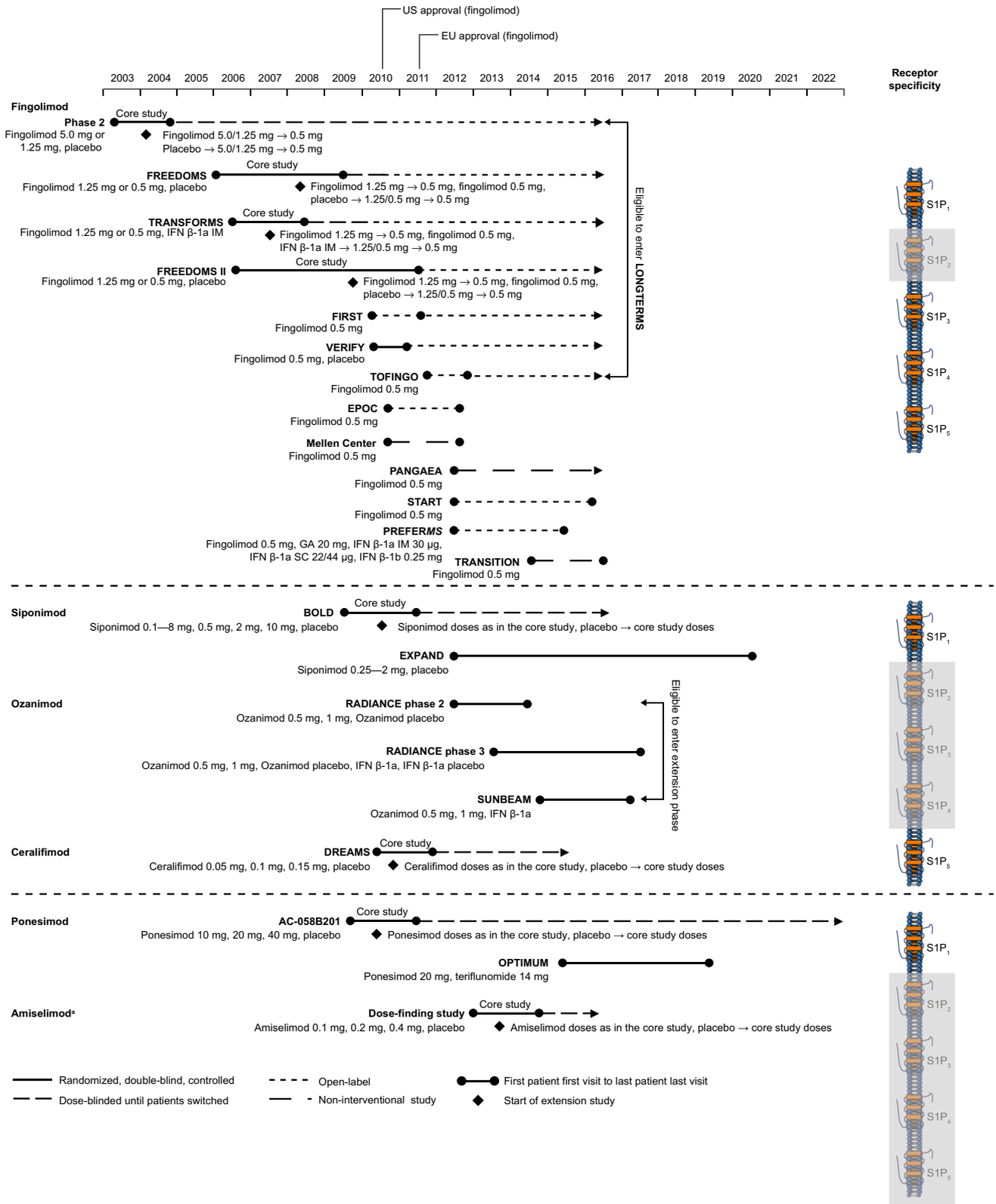
In the TRANSFORMS extension, the safety profile was consistent with that observed in the core study [46]. The incidences of AEs (94.7% and 92.2%) and SAEs (15.4 and 12.6%) were similar in the continuous-fingolimod and IFN $\beta$ -1a-fingolimod switch groups, respectively [46].

Transient effects on heart rate at treatment initiation and effects on liver function were observed, but there were no cases of macular edema [46]. The incidence of herpes viral infections was similar in both treatment groups (continuous-fingolimod group,  $n = 36$  [10.1%]; treatment-switch group,  $n = 25$  [15%]) [46]. There was an imbalance in the incidence of cancer between the continuous-fingolimod and treatment-switch groups ( $n = 14$  and  $n = 1$ , respectively) [46], but the incidences of basal cell carcinoma and other cancer types were similar to those reported in a pooled analysis of 3553 patients treated with fingolimod in the core and extension trials [46].

In the LONGTERMS study, the safety profile of fingolimod was confirmed at interim analysis, and no long-term safety or tolerability concerns were raised [52, 53]. Incidence rate ratios [95% confidence interval (CI)] for AEs and SAEs were 0.74 (0.68–0.79) and 0.73 (0.60–0.91), respectively, relative to events reported in a cohort of patients pooled from the phase III trials, suggesting that incidence does not increase with prolonged fingolimod exposure [52, 53]. Incidence rate ratios for AEs of special interest (including bradyarrhythmia post first-dose, hypertension, infections, liver transaminase elevation, lymphopenia, macular edema, malignant neoplasms, reproductive toxicity, respiratory conditions, skin cancer, and thromboembolic events) were  $<1.0$ , with the exception of lymphopenia [52, 53]. Physicians had access to lymphocyte counts in LONGTERMS, but these data were blinded in the phase III trials unless they reached a threshold of  $0.2 \times 10^9/L$ ; therefore, lymphopenia may have been reported more frequently in LONGTERMS [52, 53]. The incidence of lymphopenia was not associated with an increased risk of infection (incidence rate ratio, 0.7), and no opportunistic infections occurred in patients with sustained low actual lymphocyte counts [52]. Results of a 36-month interim analysis [49] of the PANGAEA study [50] also supported the positive safety and tolerability profiles seen in the phase III trials [49]. In the first and second years, 10.8 and 9.1% of patients discontinued therapy, 4.3 and 2.3% owing to AEs, respectively; 45.4% of patients experienced no AEs [49].

## 7 S1P Receptor Modulator Switch Studies

Studies examining switching treatment to an S1P receptor modulator are currently confined to studies of fingolimod. In the randomized 12-month PREFERMS study of treatment retention with fingolimod and injectable DMTs (iDMTs), retention was significantly greater with fingolimod (81.3% [ $n = 352$ ] vs. 29.2% [ $n = 125$ ];  $p < 0.0001$ , respectively). Among patients who switched treatment, fewer discontinued fingolimod [1.1% ( $n = 5$ )]



◀ **Fig. 2** Study design of the pivotal and extension studies of fingolimod, and of the other sphingosine-1-phosphate (S1P) receptor modulators. Areas shaded gray indicate that the S1P receptor subtypes are unaffected by the drugs in each section. <sup>a</sup>Amiselimod is selective, but its selectivity is unknown. **BOLD** BAF312 on MRI lesion given once daily, **DREAMS** a study of the safety and efficacy of ONO-4641 in patients with relapsing–remitting multiple sclerosis, **EPOC** evaluate patient outcomes, **EU** European Union, **EXPAND** exploring the efficacy and safety of siponimod in patients with secondary progressive multiple sclerosis, **FIRST** fingolimod initiation and cardiac safety trial, **FREEDOMS** FTY720 research evaluating effects of daily oral therapy in multiple sclerosis, **GA** glatiramer acetate, **IFN** interferon, **IM** intramuscular, **LONGTERMS** long-term study of fingolimod in multiple sclerosis patients from the FTY clinical program, **MOMENTUM** a randomized, double-blind, placebo-controlled, phase II, dose-finding study of MT-1303 in subjects with relapsing–remitting multiple sclerosis, **OPTIMUM** oral ponesimod versus teriflunomide in relapsing multiple sclerosis, **PANGAEA** post authorization noninterventional German safety of Gilenya in relapsing–remitting multiple sclerosis patients, **PREFERMS** evaluation of patient retention of fingolimod versus currently approved disease-modifying therapy in patients with relapsing–remitting multiple sclerosis, **RADIANCE** safety and efficacy of the selective sphingosine-1-phosphate receptor modulator ozanimod in relapsing multiple sclerosis, **SIP<sub>1-5</sub>** sphingosine-1-phosphate receptor subtypes 1–5, **SC** subcutaneous, **START** study to validate telemetric ECG systems for first-dose administration of fingolimod, **SUNBEAM** phase III study of RPC1063 in relapsing multiple sclerosis, **TOFINGO** disease control and safety in patients with relapsing–remitting multiple sclerosis switching from natalizumab to fingolimod, **TRANSFORMS** trial assessing injectable interferon versus FTY720 oral in relapsing–remitting multiple sclerosis, **TRANSITION** a two-year observational study to evaluate the safety profile of fingolimod in patients with multiple sclerosis who switch from natalizumab to fingolimod, **VERIFY** investigating the effect of recent immunization in patients receiving fingolimod therapy. Adapted with permission from Khatri, *Ther Adv Neurol Disord.* 2016;9:130–47 [47]

than iDMTs [9.8% ( $n = 43$ )], and patients receiving fingolimod also experienced greater treatment satisfaction at all assessments than those given iDMT ( $p < 0.001$ ). The rate of AEs per patient-year was lower with fingolimod (4.01) than with iDMTs (7.01) [39].

TRANSITION is a 2-year, prospective, observational, single-cohort, real-world study examining the safety profile of fingolimod in patients with RMS who switched from natalizumab (target enrollment:  $N = 1500$ ) [54]. In the 1-year interim analysis ( $n = 639$ ) [55], AEs and SAEs were experienced by 48 and 8% of patients, respectively [54]. Among AEs of interest, exposure-adjusted incidences (95% CI) per 100 patient-years were: hypertension, 2.80 (1.66–4.42); lymphopenia, 1.85 (0.96–3.23); leukopenia, 0.92 (0.34–1.99); liver transaminase elevation, 0.46 (0.09–1.34); second-degree AV block, 0.30 (0.04–1.10); macular edema, 0.30 (0.04–1.10); basal cell carcinoma, 0.30 (0.04–1.10); and infections, 15.2 (12.2–18.7). The incidence of infections reported as SAEs was 1.38 (0.63–2.62) [55].

Evaluate patient outcomes (EPOC) was a randomized, open-label, 6-month, multicenter study of treatment

satisfaction in patients who switched from an iDMT to fingolimod 0.5 mg without a washout period, compared with patients who remained on iDMT or switched to an alternative iDMT [56]. Patient-reported satisfaction measures (including effectiveness, side effects, and convenience), and measures of fatigue and depression, were significantly improved in patients who switched to fingolimod compared with those receiving iDMT (all  $p < 0.001$ ) [56]. Adverse events and SAEs were experienced by 78.8 and 4% of patients taking fingolimod, and by 62 and 2% of patients on iDMTs, respectively. The three most frequent AEs in the fingolimod group were headache (12.4%), fatigue (11.5%), and upper respiratory tract infection (6.5%). Multiple sclerosis relapse (0.6%), lymphopenia (0.3%), and non-cardiac chest pain (0.3%) were the only SAEs with more than one case in the fingolimod group, while only MS relapse was reported in the iDMT group (0.8%). Adverse events leading to study-drug discontinuation occurred in 5.2% of patients taking fingolimod (most commonly macular edema [0.5%] and fatigue [0.5%]) compared with 1.6% of patients taking iDMT. Infections were reported in similar proportions of patients in the fingolimod group (30.1%) and the iDMT group (27.8%). Similarly to LONGTERMS, low lymphocyte counts were not associated with increased infection rate in fingolimod-treated patients, and no significant differences were found in lymphocyte counts between fingolimod-treated patients who experienced an infection and those who did not [56].

The randomized, double-blind, placebo-controlled, 32-week TOFINGO study examined the effect of washout duration in patients with relapsing–remitting multiple sclerosis switching from natalizumab to fingolimod ( $n = 142$ ) [57]. Adverse events were experienced by 68% of patients and were mostly mild or moderate with a similar incidence across groups. No unusually severe relapses or opportunistic infections occurred after treatment switch, and shorter washout periods were associated with less MRI disease activity than longer periods [57].

## 8 Safety Precautions for Fingolimod Therapy

As the only approved S1P receptor modulator, fingolimod has the greatest amount of safety data available, together with a large amount of clinical experience, with total patient exposure exceeding 397,000 patient-years [37]. Factors that should be considered before initiation of fingolimod or that require monitoring on treatment, including first-dose monitoring, pregnancy, diabetes mellitus, posterior reversible encephalopathy syndrome, basal cell carcinoma, infections such as varicella, opportunistic cryptococcal infections, and progressive multifocal

**Table 2** Safety precautions before and during fingolimod therapy

Safety consideration	Recommendation
First-dose monitoring [6, 7, 58]	Vital signs, such as blood pressure and heart rate, and electrocardiogram assessments should be recorded before administering fingolimod, and patients should be monitored for at least 6 h after their first dose, or until heart rate has passed its nadir, and they are asymptomatic for reduced heart rate. Most cardiovascular effects resolve without intervention
Pregnancy [6, 7]	When applicable, a negative pregnancy test should be obtained before initiation of fingolimod, and the need for effective contraception during treatment discussed. If a woman becomes pregnant while taking fingolimod, treatment discontinuation is recommended. Furthermore, because it takes approximately 2 months to eliminate fingolimod from the body, pregnancy should be avoided for 2 months after stopping treatment
Macular edema [6, 7]	Examination of the fundus including the macula is required in all patients before starting treatment, at 3–4 months after starting treatment, and again at any time after a patient reports visual disturbances while receiving fingolimod therapy
Diabetes mellitus [6, 7, 59]	Fingolimod should be used with caution in patients with diabetes owing to a potential increase in the risk of macular edema. Regular ophthalmological examinations are advised to detect macular edema. Incidence of macular edema (0.3%) appears low, and macular changes do not progress after discontinuation
Posterior reversible encephalopathy syndrome [6, 7]	Rare cases of posterior reversible encephalopathy syndrome have been reported, and if suspected, fingolimod should be discontinued
Basal cell carcinoma [6, 7]	Basal cell carcinoma has also been reported in patients receiving fingolimod. Vigilance for suspicious skin lesions is warranted, and patients should be referred to a dermatologist if lesions are detected
Infections, such as varicella (chickenpox) [6, 7, 58–60]	Before initiating treatment with fingolimod, patients' immunity to varicella (chickenpox) should be assessed, and they should be vaccinated if they are seronegative. Initiation of fingolimod therapy should be postponed in patients with severe active infections until all infections are resolved. A safety analysis of over 3400 patients taking fingolimod indicated no elevated risk of infectious adverse events compared with placebo
Opportunistic cryptococcal infections [6, 7, 58–60]	Cases of opportunistic cryptococcal infections have been reported rarely for patients receiving fingolimod. When such infections have occurred, they have typically been in patients with over 2 years of exposure to fingolimod. Patients with symptoms consistent with a cryptococcal infection should be evaluated and treated immediately
PML [6, 7, 58–61]	Fingolimod is classified as a disease-modifying therapy with a low risk of PML. Cases of PML have been reported rarely for patients receiving fingolimod. When such infections have occurred, they have typically been in patients with over 2 years of exposure to fingolimod. Vigilance should be maintained for signs of PML, and patients with symptoms suggestive of PML should be evaluated and treated immediately

*PML* progressive multifocal leukoencephalopathy

leukoencephalopathy, are summarized in Table 2 [6, 7, 58–61].

First-dose monitoring for cardiovascular effects is required with fingolimod treatment and is likely to be required for the second generation of S1P receptor modulators; and as such is unlikely to be discontinued in the near future. However, clinical data with fingolimod indicate that the first-dose effects are not as widespread or as severe as previously thought, allowing the first dose of fingolimod treatment to be initiated outside of the clinic setting. Interim results from a study of fingolimod initiation ( $n = 3951$ ) reported that 0.7% of patients experienced a bradycardia of  $<45$  bpm, and 1.6% had an AV block of grade  $\geq 2$  following the first dose [58]. Furthermore, limited first-dose effects have been observed with ozanimod and amiselimod [28, 36]; therefore, the need for first-dose monitoring may be reduced in future.

Fingolimod showed developmental toxicity in animal studies, and the S1P receptor is known to be involved in vascular formation during embryogenesis; women should therefore avoid becoming pregnant while taking fingolimod, and treatment discontinuation is recommended if pregnancy occurs [6, 7]. Minor dose-dependent reductions in values for forced expiratory volume and diffusion capacity for carbon monoxide were observed with fingolimod treatment, starting as early as 1 month after treatment initiation and remaining stable thereafter [6, 7]. Fingolimod should therefore be used with caution in patients with severe respiratory disease, pulmonary fibrosis, or chronic obstructive pulmonary disease [6]. Basal cell carcinoma has been associated with fingolimod use, with increased incidence reported in clinical trials compared with placebo; healthcare professionals and patients are therefore advised to monitor for suspicious skin lesions

[6, 7]. Tumefactive lesions, which can be part of a spectrum of the MS course itself, have been noted with fingolimod during therapy as well as after withdrawal of fingolimod [62–66]. As with natalizumab, a high-efficacy DMT, disease reactivation, often labeled as ‘rebound’, has been reported after withdrawal of fingolimod therapy [67–71].

A dose-dependent increase in the risk of macular edema was observed in the fingolimod clinical development program [6, 7]. Macular edema occurred predominantly during the first 3–4 months of therapy and generally resolved after fingolimod discontinuation; macular edema has also been reported in the post-marketing setting, usually within the first 6 months of therapy. Patients are required to undergo examination of the fundus including the macula prior to treatment and at 3–4 months after starting treatment, and should also be examined at any time when they report visual disturbances while receiving therapy [6, 7]. Patients with a history of uveitis and patients with diabetes mellitus are at increased risk of macular edema with fingolimod; thus regular follow-up examinations are recommended for these patients during treatment [6, 7].

The dose-dependent reduction in peripheral lymphocyte count with fingolimod treatment can increase the risk of infections [6, 7, 60]. In clinical trials, some infections (such as herpes virus and respiratory tract infections) were more common with fingolimod than with placebo, and there was a higher incidence of serious infections with fingolimod than with placebo. Serious infections with opportunistic pathogens including viruses, fungi, and bacteria have been reported in the post-marketing setting [6, 7]. These include serious life-threatening events of disseminated varicella zoster and herpes simplex infections (including one fatal event), and cases of Kaposi’s sarcoma. Cryptococcal infections have also been reported, albeit rarely, with fingolimod in the post-marketing setting, typically occurring after approximately 2–3 years of treatment [6, 7, 60]. Patients with signs or symptoms consistent with any of these infections should undergo prompt diagnostic evaluation and appropriate management.

Progressive multifocal leukoencephalopathy (PML) is an opportunistic viral infection of the brain caused by the John Cunningham virus; it typically occurs only in patients who are immunocompromised, and may be fatal or result in severe disability. Cases of PML have been reported in patients treated with fingolimod [6, 7, 61]. In many instances, patients had received prior treatment with natalizumab, which is associated with an increased risk of PML [61]. However, as of February 2017, 13 cases of probable or definite PML have been reported in patients treated with fingolimod without prior natalizumab therapy [61, 72]. These cases occurred in patients who had been receiving fingolimod therapy for between 18 and

65 months, and who were aged 34–63 years [61, 72]. These cases of PML were not associated with sustained grade 4 lymphopenia [61]; more research is needed to determine the relationship between lymphopenia and the risk of developing PML [73]. The low frequency of PML with fingolimod therapy means that there is currently limited information available to identify factors associated with an increased risk of this infection. Fingolimod should be withheld at the first sign or symptom suggestive of PML and an appropriate diagnostic evaluation performed; vigilance for PML-suspicious lesions during routine MRI is also important [6, 7]. In a recent article contextualizing the risk of PML with DMTs in MS, natalizumab was classified as having a high potential risk of PML, while fingolimod was grouped together with dimethyl fumarate as having a low risk [61, 72].

## 9 Conclusions

The ability of S1P receptor modulators to block immune cell trafficking specifically [3, 8, 11, 15–17] makes them particularly interesting for the treatment of a number of autoimmune diseases including MS [1, 3]. Fingolimod is the only approved agent with extensively characterized efficacy and safety profiles, but several S1P receptor modulators are currently in development. The specificity of the immunomodulatory effects of S1P receptor modulators suggests that the likelihood of off-target AEs in the long term is small, unlike with other drugs licensed for the treatment of MS associated with broad-range immune suppression. The short-term efficacy data available for the S1P receptor modulators under development support the potential effectiveness of this drug class in MS, and the available safety data are consistent across the class and with the known mechanism of action. The long-term safety, tolerability, and persistence profiles of fingolimod, including upon switching from other treatments, should reassure clinicians regarding the suitability of S1P receptor modulators as treatment options for patients with MS.

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