ANNEX I

SUMMARY OF PRODUCT CHARACTERISTICS
1. NAME OF THE MEDICINAL PRODUCT

Tecfidera 120 mg gastro-resistant hard capsules

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each capsule contains 120 mg dimethyl fumarate.

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Gastro-resistant hard capsule

Green and white gastro-resistant hard capsule printed with ‘BG-12 120 mg’.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Tecfidera is indicated for the treatment of adult patients with relapsing remitting multiple sclerosis (please refer to section 5.1 for important information on the populations for which efficacy has been established).

4.2 Posology and method of administration

Treatment should be initiated under supervision of a physician experienced in the treatment of the disease.

Posology

The starting dose is 120 mg twice a day. After 7 days, the dose is increased to the recommended dose of 240 mg twice a day.

Temporary dose reduction to 120 mg twice a day may reduce the occurrence of flushing and gastrointestinal adverse reactions. Within 1 month, the recommended dose of 240 mg twice a day should be resumed.

Tecfidera should be taken with food (see section 5.2). For those patients who may experience flushing or gastrointestinal adverse reactions, taking Tecfidera with food may improve tolerability (see sections 4.4, 4.5 and 4.8).

Older people

Clinical studies of Tecfidera had limited exposure to patients aged 55 years and above, and did not include sufficient numbers of patients aged 65 and over to determine whether they respond differently than younger patients (see section 5.2). Based on the mode of action of the active substance there are no theoretical reasons for any requirement for dose adjustments in the elderly.
Renal and hepatic impairment

Tecfidera has not been studied in patients with renal or hepatic impairment. Based on clinical pharmacology studies, no dose adjustments are needed (see section 5.2). Caution should be used when treating patients with severe renal or severe hepatic impairment (see section 4.4).

Paediatric population

The safety and efficacy of Tecfidera in children and adolescents aged 10 to 18 years have not been established. No data are available. There is no relevant use of Tecfidera in children aged less than 10 years in multiple sclerosis.

Method of administration

For oral use.

The capsule or its contents should not be crushed, divided, dissolved, sucked or chewed as the enteric-coating of the microtablets prevents irritant effects on the gut.

4.3 Contraindications

Hypersensitivity to the active substance or to any of the excipients listed in section 6.1.

4.4 Special warnings and precautions for use

Blood/laboratory tests

Tecfidera may decrease lymphocyte counts (see section 4.8). Tecfidera has not been studied in patients with pre-existing low lymphocyte counts and caution should be exercised when treating these patients. Prior to initiating treatment with Tecfidera, a recent complete blood count (i.e. within 6 months) should be available. Assessments of complete blood counts are also recommended after 6 months of treatment and every 6 to 12 months thereafter and as clinically indicated.

Changes in renal and hepatic laboratory tests have been seen in clinical trials in subjects treated with Tecfidera (see section 4.8). The clinical implications of these changes are unknown. Assessments of renal function (e.g. creatinine, blood urea nitrogen and urinalysis) and hepatic function (e.g. ALT and AST) are recommended prior to treatment initiation, after 3 and 6 months of treatment, every 6 to 12 months thereafter and as clinically indicated.

Severe renal and hepatic impairment

Tecfidera has not been studied in patients with severe renal or severe hepatic impairment and caution should, therefore, be used in these patients (see section 4.2).

Severe active gastrointestinal disease

Tecfidera has not been studied in patients with severe active gastrointestinal disease and caution should, therefore, be used in these patients.

Flushing

In clinical trials, 34% of Tecfidera treated patients experienced flushing. In the majority of patients who experienced flushing, it was mild or moderate in severity.

In clinical trials, 3 patients out of a total of 2,560 patients treated with Tecfidera experienced serious flushing symptoms that were probable hypersensitivity or anaphylactoid reactions. These events were not life-threatening, but led to hospitalisation. Prescribers and patients should be alert to this
possibility in the event of severe flushing reactions (see sections 4.2, 4.5 and 4.8).

Infections

In phase III placebo-controlled studies, the incidence of infections (60% vs 58%) and serious infections (2% vs 2%) was similar in patients treated with Tecfidera or placebo, respectively. There was no increased incidence of serious infections observed in patients with lymphocyte counts <0.8x10^9/L or <0.5x10^9/L. During treatment with Tecfidera in the MS placebo controlled trials, mean lymphocyte counts decreased by approximately 30% from baseline at one year and then plateaued (see section 4.8). Mean lymphocyte counts remained within normal limits. If a patient develops a serious infection, suspending treatment with Tecfidera should be considered and the benefits and risks should be reassessed prior to re-initiation of therapy. Patients receiving Tecfidera should be instructed to report symptoms of infections to a physician. Patients with serious infections should not start treatment with Tecfidera until the infection(s) is resolved.

4.5 Interaction with other medicinal products and other forms of interaction

Tecfidera has not been studied in combination with anti-neoplastic or immunosuppressive therapies and caution should, therefore, be used during concomitant administration. In multiple sclerosis clinical studies, the concomitant treatment of relapses with a short course of intravenous corticosteroids was not associated with a clinically relevant increase of infection.

Vaccination during treatment with Tecfidera has not been studied. It is not known whether treatment with Tecfidera might reduce the effectiveness of some vaccines. Live vaccines might carry an increased risk of clinical infection and should not be given to patients treated with Tecfidera unless, in exceptional cases, this potential risk is considered to be outweighed by the risk to the individual of not vaccinating.

During treatment with Tecfidera, simultaneous use of other fumaric acid derivatives (topical or systemic) should be avoided.

In humans, dimethyl fumarate is extensively metabolised by esterases before it reaches the systemic circulation and further metabolism occurs through the tricarboxylic acid cycle, with no involvement of the cytochrome P450 (CYP) system. Potential drug interaction risks were not identified from in vitro CYP-inhibition and induction studies, a p-glycoprotein study, or studies of the protein binding of dimethyl fumarate and monomethyl fumarate (a primary metabolite of dimethyl fumarate).

Commonly used medicinal products in patients with multiple sclerosis, intramuscular interferon beta-1a and glatiramer acetate, were clinically tested for potential interactions with dimethyl fumarate and did not alter the pharmacokinetic profile of dimethyl fumarate.

Administration of 325 mg (or equivalent) non-enteric coated acetylsalicylic acid, 30 minutes prior to Tecfidera, over 4 days of dosing, did not alter the pharmacokinetic profile of Tecfidera and reduced the occurrence and severity of flushing in a healthy volunteer study. However, long term use of acetylsalicylic acid is not recommended for the management of flushing. Potential risks associated with acetylsalicylic acid therapy should be considered prior to co-administration with Tecfidera. (see sections 4.2, 4.4 and 4.8).

Concurrent therapy with nephrotoxic medicinal products (such as aminoglycosides, diuretics, NSAIDs or lithium) may increase the potential of renal adverse reactions (e.g. proteinuria) in patients taking Tecfidera (see section 4.8).

Consumption of moderate amounts of alcohol did not alter exposure to Tecfidera and was not associated with an increase in adverse reactions. Consumption of large quantities of undiluted strong alcoholic drinks (more than 30% alcohol by volume) may lead to increased dissolution rates of Tecfidera and, therefore, may increase the frequency of gastrointestinal adverse reactions.
In vitro CYP induction studies did not demonstrate an interaction between Tecfidera and oral contraceptives. In vivo interaction studies have not been performed with oral contraceptives. Even though an interaction is not expected, non-hormonal contraceptive measures should be considered with Tecfidera (see section 4.6).

Paediatric population

Interaction studies have only been performed in adults.

4.6 Fertility, pregnancy and lactation

Pregnancy

There are no or limited amount of data from the use of dimethyl fumarate in pregnant women. Animal studies have shown reproductive toxicity (see section 5.3). Tecfidera is not recommended during pregnancy and in women of childbearing potential not using appropriate contraception (see section 4.5). Tecfidera should be used during pregnancy only if clearly needed and if the potential benefit justifies the potential risk to the foetus.

Breast-feeding

It is unknown whether dimethyl fumarate or its metabolites are excreted in human milk. A risk to the newborns/infants cannot be excluded. A decision must be made whether to discontinue breast-feeding or to discontinue Tecfidera therapy. The benefit of breast-feeding for the child and the benefit of therapy for the woman should be taken into account.

Fertility

There are no data on the effects of Tecfidera on human fertility. Data from preclinical studies do not suggest that dimethyl fumarate would be associated with an increased risk of reduced fertility (see section 5.3).

4.7 Effects on ability to drive and use machines

No studies on the ability to drive and use machines have been conducted.

4.8 Undesirable effects

Summary of the safety profile

The most common adverse reactions (incidence ≥10%) for patients treated with Tecfidera were flushing and gastrointestinal events (i.e. diarrhoea, nausea, abdominal pain, abdominal pain upper). Flushing and gastrointestinal events tend to begin early in the course of treatment (primarily during the first month) and in patients who experience flushing and gastrointestinal events, these events may continue to occur intermittently throughout treatment with Tecfidera. The most commonly reported adverse reactions leading to discontinuation (incidence >1%) in patients treated with Tecfidera were flushing (3%) and gastrointestinal events (4%).

In placebo-controlled and uncontrolled clinical studies, a total of 2,468 patients have received Tecfidera and been followed for periods up to 4 years with an overall exposure equivalent to 3,588 person-years. Approximately 1,056 patients have received more than 2 years of treatment with Tecfidera. The experience in uncontrolled clinical trials is consistent with the experience in the placebo-controlled clinical trials.
Tabulated summary of adverse reactions

Adverse reactions, which were more frequently reported in Tecfidera versus placebo-treated patients, are presented in the table below. These data were derived from 2 pivotal Phase 3 placebo-controlled, double-blind clinical trials with a total of 1,529 patients treated with Tecfidera and for up to 24 months with an overall exposure of 2,371 person-years (see section 5.1). The frequencies described in the table below are based on 769 patients treated with Tecfidera 240 mg twice a day and 771 patients treated with placebo.

The adverse reactions are presented as MedDRA preferred terms under the MedDRA System Organ Class. The incidence of the adverse reactions below are expressed according to the following categories:
- Very common (≥1/10)
- Common (≥1/100 to <1/10)
- Uncommon (≥1/1,000 to <1/100)
- Rare (≥1/10,000 to <1/1,000)
- Very rare (<1/10,000)
- Not known (cannot be estimated from the available data)

<table>
<thead>
<tr>
<th>MedDRA System Organ Class</th>
<th>Adverse reaction</th>
<th>Frequency category</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infections and infestations</td>
<td>Gastroenteritis</td>
<td>Common</td>
</tr>
<tr>
<td>Blood and lymphatic system disorders</td>
<td>Lymphopenia</td>
<td>Common</td>
</tr>
<tr>
<td></td>
<td>Leucopenia</td>
<td>Common</td>
</tr>
<tr>
<td>Immune system disorders</td>
<td>Hypersensitivity</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Nervous system disorders</td>
<td>Burning sensation</td>
<td>Common</td>
</tr>
<tr>
<td>Vascular disorders</td>
<td>Flushing</td>
<td>Very common</td>
</tr>
<tr>
<td></td>
<td>Hot flush</td>
<td>Common</td>
</tr>
<tr>
<td>Gastrointestinal disorders</td>
<td>Diarrhoea</td>
<td>Very common</td>
</tr>
<tr>
<td></td>
<td>Nausea</td>
<td>Very common</td>
</tr>
<tr>
<td></td>
<td>Abdominal pain upper</td>
<td>Very common</td>
</tr>
<tr>
<td></td>
<td>Abdominal pain</td>
<td>Very common</td>
</tr>
<tr>
<td></td>
<td>Vomiting</td>
<td>Common</td>
</tr>
<tr>
<td></td>
<td>Dyspepsia</td>
<td>Common</td>
</tr>
<tr>
<td></td>
<td>Gastritis</td>
<td>Common</td>
</tr>
<tr>
<td></td>
<td>Gastrointestinal disorder</td>
<td>Common</td>
</tr>
<tr>
<td>Skin and subcutaneous tissue disorders</td>
<td>Pruritus</td>
<td>Common</td>
</tr>
<tr>
<td></td>
<td>Rash</td>
<td>Common</td>
</tr>
<tr>
<td></td>
<td>Erythema</td>
<td>Common</td>
</tr>
<tr>
<td>Renal and urinary disorders</td>
<td>Proteinuria</td>
<td>Common</td>
</tr>
<tr>
<td>General disorders and administration site conditions</td>
<td>Feeling hot</td>
<td>Common</td>
</tr>
<tr>
<td>Investigations</td>
<td>Ketones measured in urine</td>
<td>Very common</td>
</tr>
<tr>
<td></td>
<td>Albumin urine present</td>
<td>Common</td>
</tr>
<tr>
<td></td>
<td>Aspartate aminotransferase increased</td>
<td>Common</td>
</tr>
<tr>
<td></td>
<td>Alanine aminotransferase increased</td>
<td>Common</td>
</tr>
<tr>
<td></td>
<td>White blood cell count decreased</td>
<td>Common</td>
</tr>
</tbody>
</table>

Description of selected adverse reactions

**Flushing**

In the placebo-controlled studies, the incidence of flushing (34% versus 4%) and hot flush (7% versus 2%) was increased in patients treated with Tecfidera compared to placebo, respectively. Flushing is usually described as flushing or hot flush, but can include other events (e.g. warmth, redness, itching, and burning sensation). Flushing events tend to begin early in the course of treatment (primarily during the first month) and in patients who experience flushing, these events may continue
to occur intermittently throughout treatment with Tecfidera. In patients with flushing, the majority had flushing events that were mild or moderate in severity. Overall, 3% of patients treated with Tecfidera discontinued due to flushing. The incidence of serious flushing, which may be characterised by generalised erythema, rash and/or pruritus, was seen in less than 1% of patients treated with Tecfidera (see sections 4.2, 4.4 and 4.5).

**Gastrointestinal**

The incidence of gastrointestinal events (e.g. diarrhoea [14% versus 10%], nausea [12% versus 9%], upper abdominal pain [10% versus 6%], abdominal pain [9% versus 4%], vomiting [8% versus 5%] and dyspepsia [5% versus 3%]) was increased in patients treated with Tecfidera compared to placebo, respectively. Gastrointestinal events tend to begin early in the course of treatment (primarily during the first month) and in patients who experience gastrointestinal events, these events may continue to occur intermittently throughout treatment with Tecfidera. In the majority of patients who experienced gastrointestinal events, it was mild or moderate in severity. Four per cent (4%) of patients treated with Tecfidera discontinued due to gastrointestinal events. The incidence of serious gastrointestinal events, including gastroenteritis and gastritis, was seen in 1% of patients treated with Tecfidera (see section 4.2).

**Hepatic transaminases**

In placebo-controlled studies, elevations of hepatic transaminases were observed. The majority of patients with elevations had hepatic transaminases that were <3 times the upper limit of normal (ULN). The increased incidence of elevations of hepatic transaminases in patients treated with Tecfidera relative to placebo was primarily seen during the first 6 months of treatment. Elevations of alanine aminotransferase and aspartate aminotransferase ≥3 times ULN, respectively, were seen in 5% and 2% of patients treated with placebo and 6% and 2% of patients treated with Tecfidera. There were no elevations in transaminases ≥3 times ULN with concomitant elevations in total bilirubin >2 times ULN. Discontinuations due to elevated hepatic transaminases were <1% and similar in patients treated with Tecfidera or placebo.

**Renal**

In placebo-controlled studies, the incidence of proteinuria was higher in patients treated with Tecfidera (9%) compared to placebo (7%). The overall incidence of renal and urinary adverse events was similar for Tecfidera and placebo-treated patients. There were no reports of serious renal failure. On urinalysis, the percentage of patients with protein values of 1+ or greater was similar for Tecfidera (43%) and placebo-treated patients (40%). Typically, laboratory observations of proteinuria were not progressive. Compared to patients treated with placebo, estimated glomerular filtration rate (eGFR) was observed to increase in patients treated with Tecfidera, including those patients with 2 consecutive occurrences of proteinuria (≥1+).

**Haematological**

In the placebo-controlled studies, most patients (>98%) had normal lymphocyte values prior to initiating treatment. Upon treatment with Tecfidera, mean lymphocyte counts decreased over the first year with a subsequent plateau. On average, lymphocyte counts decreased by approximately 30% of baseline value. Mean and median lymphocyte counts remained within normal limits. Lymphocyte counts <0.5x10⁹/l were observed in <1% of patients treated with placebo and 6% of patients treated with Tecfidera. A lymphocyte count <0.2x10⁹/l was observed in 1 patient treated with Tecfidera and in no patients treated with placebo. The incidence of infections (58% versus 60%) and serious infections (2% versus 2%) was similar in patients treated with placebo or Tecfidera. An increased incidence of infections and serious infections was not observed in patients with lymphocyte counts <0.8x10⁹/l or <0.5x10⁹/l. A transient increase in mean eosinophil counts was seen during the first 2 months of therapy.
Laboratory abnormalities

In the placebo-controlled studies, measurement of urinary ketones (1+ or greater) was higher in patients treated with Tecfidera (45%) compared to placebo (10%). No untoward clinical consequences were observed in clinical trials.

Levels of 1,25-dihydroxyvitamin D decreased in Tecfidera treated patients relative to placebo (median percentage decrease from baseline at 2 years of 25% versus 15%, respectively) and levels of parathyroid hormone (PTH) increased in Tecfidera treated patients relative to placebo (median percentage increase from baseline at 2 years of 29% versus 15%, respectively). Mean values for both parameters remained within normal range.

Reporting of suspected adverse reactions

Reporting suspected adverse reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/risk balance of the medicinal product. Healthcare professionals are asked to report any suspected adverse reactions via the national reporting system listed in Appendix V.

4.9 Overdose

No cases of overdose have been reported.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Other nervous system drugs, ATC code: N07XX09

Mechanism of action

The mechanism by which dimethyl fumarate exerts therapeutic effects in multiple sclerosis is not fully understood. Preclinical studies indicate that dimethyl fumarate pharmacodynamic responses appear to be primarily mediated through activation of the Nuclear factor (erythroid-derived 2)-like 2 (Nrf2) transcriptional pathway. Dimethyl fumarate has been shown to up regulate Nrf2-dependent antioxidant genes in patients (e.g. NAD(P)H dehydrogenase, quinone 1; [NQO1]).

Pharmacodynamic effects

Effects on the immune system

In preclinical and clinical studies, Tecfidera demonstrated anti-inflammatory and immunomodulatory properties. Dimethyl fumarate and monomethyl fumarate, the primary metabolite of dimethyl fumarate, significantly reduced immune cell activation and subsequent release of pro-inflammatory cytokines in response to inflammatory stimuli in preclinical models. In clinical studies with psoriasis patients, dimethyl fumarate affected lymphocyte phenotypes through a down-regulation of pro-inflammatory cytokine profiles (T_{h1}, T_{h17}), and biased towards anti-inflammatory production (T_{h2}). Dimethyl fumarate demonstrated therapeutic activity in multiple models of inflammatory and neuroinflammatory injury. In Phase 3 studies, upon treatment with Tecfidera mean lymphocyte counts decreased on average by approximately 30% of their baseline value over the first year with a subsequent plateau.

Effect on cardiovascular system

Single doses of 240 mg or 360 mg Tecfidera did not have any effect on the QTc interval when compared to placebo in a QTc study.
Clinical efficacy and safety

Two, 2-year, randomised, double-blind, placebo controlled studies [Study 1 (DEFINE) with 1234 subjects and Study 2 (CONFIRM) with 1417 subjects] of subjects with relapsing-remitting multiple sclerosis (RRMS) were performed. Subjects with progressive forms of MS were not included in these studies. Efficacy (see table below) and safety were demonstrated in subjects with Expanded Disability Status Scale (EDSS) scores ranging from 0 to 5 inclusive, who had experienced at least 1 relapse during the year prior to randomisation, or, within 6 weeks of randomisation had a brain Magnetic Resonance Imaging (MRI) demonstrating at least one gadolinium-enhancing (Gd+) lesion. Study 2 contained a rater-blinded (i.e. study physician/ investigator assessing the response to study treatment was blinded) reference comparator of glatiramer acetate.

In Study 1, patients had the following median baseline characteristics: age 39 years, disease duration 7.0 years, EDSS score 2.0. In addition, 16% of patients had an EDSS score >3.5, 28% had ≥2 relapses in the prior year and 42% had previously received other approved MS treatments. In the MRI cohort 36% of patients entering the study had Gd+ lesions at baseline (mean number of Gd+ lesions 1.4).

In Study 2, patients had the following median baseline characteristics: age 37 years, disease duration 6.0 years, EDSS score 2.5. In addition, 17% of patients had an EDSS score >3.5, 32% had ≥2 relapses in the prior year and 30% had previously received other approved MS treatments. In the MRI cohort 45% of patients entering the study had Gd+ lesions at baseline (mean number of Gd+ lesions 2.4).

Compared to placebo, subjects treated with Tecfidera had a clinically meaningful and statistically significant reduction on: the primary endpoint in Study 1, proportion of subjects relapsed at 2 years; and the primary endpoint in Study 2, annualised relapse rate at 2 years.

The annualised relapse rate for glatiramer acetate and placebo was 0.286 and 0.401 respectively in Study 2, corresponding to a reduction of 29% (p=0.013), which is consistent with approved prescribing information.

<table>
<thead>
<tr>
<th>DEFINE</th>
<th>CONFIRM</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Placebo Tecfidera Placebo Tecfidera Glatiramer 240 mg twice a day 240 mg twice a day acetate</td>
</tr>
</tbody>
</table>

| **Clinical Endpoints**
| No. subjects | Placebo | 408 | 410 | 401 | 363 | 359 | 350 |
| Annualised relapse rate | 0.364 | 0.172*** | 0.401 | 0.224*** | 0.286* |
| Rate ratio (95% CI) | 0.47 (0.37, 0.61) | 0.56 (0.42, 0.74) | 0.71 (0.55, 0.93) |
| Proportion relapsed | 0.461 | 0.270*** | 0.410 | 0.291** | 0.321** |
| Hazard ratio (95% CI) | 0.51 (0.40, 0.66) | 0.66 (0.51, 0.86) | 0.71 (0.55, 0.92) |
| Proportion with 12-week confirmed disability progression | 0.271 | 0.164** | 0.169 | 0.128* | 0.156* |
| Hazard ratio (95% CI) | 0.62 (0.44, 0.87) | 0.79 (0.52, 1.19) | 0.93 (0.63, 1.37) |
| Proportion with 24 week confirmed disability progression | 0.169 | 0.128# | 0.125 | 0.078# | 0.108# |
| Hazard ratio (95% CI) | 0.77 (0.52, 1.14) | 0.62 (0.37, 1.03) | 0.87 (0.55, 1.38) |

| **MRI Endpoints**
<p>| No. subjects | 165 | 152 | 144 | 147 | 161 |</p>
<table>
<thead>
<tr>
<th></th>
<th>DEFINE</th>
<th>CONFIRM</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Placebo</td>
<td>Tecfidera 240 mg twice a day</td>
</tr>
<tr>
<td>Mean (median) number of new or newly enlarging T2 lesions over 2 years</td>
<td>16.5 (7.0)</td>
<td>3.2 (1.0)***</td>
</tr>
<tr>
<td>Lesion mean ratio (95% CI)</td>
<td>0.15 (0.10, 0.23)</td>
<td></td>
</tr>
<tr>
<td>Mean (median) number of Gd lesions at 2 years</td>
<td>1.8 (0)</td>
<td>0.1 (0)***</td>
</tr>
<tr>
<td>Odds ratio (95% CI)</td>
<td>0.10 (0.05, 0.22)</td>
<td></td>
</tr>
<tr>
<td>Mean (median) number of new T1 hypointense lesions over 2 years</td>
<td>5.7 (2.0)</td>
<td>2.0 (1.0)***</td>
</tr>
<tr>
<td>Lesion mean ratio (95% CI)</td>
<td>0.28 (0.20, 0.39)</td>
<td></td>
</tr>
</tbody>
</table>

1 All analyses of clinical endpoints were intent-to-treat; aMRI analysis used MRI cohort *P-value < 0.05; **P-value < 0.01; ***P-value < 0.0001; #not statistically significant

Efficacy in patients with high disease activity:
Consistent treatment effect on relapses in a subgroup of patients with high disease activity was observed, whilst the effect on time to 3-month sustained disability progression was not clearly established. Due to the design of the studies, high disease activity was defined as follows:
- Patients with 2 or more relapses in one year, and with one or more Gd-enhancing lesions on brain MRI (n=42 in DEFINE; n=51 in CONFIRM) or,
- Patients who have failed to respond to a full and adequate course (at least one year of treatment) of beta-interferon, having had at least 1 relapse in the previous year while on therapy, and at least 9 T2-hyperintense lesions in cranial MRI or at least 1 Gd-enhancing lesion, or patients having an unchanged or increased relapse rate in the prior year as compared to the previous 2 years (n=177 in DEFINE; n=141 in CONFIRM).

Paediatric population

The European Medicines Agency has deferred the obligation to submit the results of studies with Tecfidera in one or more subsets of the paediatric population in multiple sclerosis (see section 4.2 for information on paediatric use).

5.2 Pharmacokinetic properties

Orally administered Tecfidera (dimethyl fumarate) undergoes rapid presystemic hydrolysis by esterases and is converted to its primary metabolite, monomethyl fumarate, which is also active. Dimethyl fumarate is not quantifiable in plasma following oral administration of Tecfidera. Therefore, all pharmacokinetic analyses related to dimethyl fumarate were performed with plasma monomethyl fumarate concentrations. Pharmacokinetic data were obtained in subjects with multiple sclerosis and healthy volunteers.

Absorption

The T_max of monomethyl fumarate is 2 to 2.5 hours. As Tecfidera gastro-resistant hard capsules contain microtablets, which are protected by an enteric coating, absorption does not commence until they leave the stomach (generally less than 1 hour). Following 240 mg twice a day administered with food, the median peak (C_max) was 1.72 mg/l and overall (AUC) exposure was 8.02 h.mg/l in subjects with multiple sclerosis. Overall, C_max and AUC increased approximately dose-proportionally in the dose range studied (120 mg to 360 mg). In subjects with multiple sclerosis, two 240 mg doses were administered 4 hours apart as part of a three times a day dosing regimen. This resulted in a minimal
accumulation of exposure yielding an increase in the median Cmax of 12% compared to the twice
daily dosing (1.72 mg/l for twice daily compared to 1.93 mg/l for three times daily) with no safety
implications.

Food does not have a clinically significant effect on exposure of dimethyl fumarate. However,
Tecfidera should be taken with food due to improved tolerability with respect to flushing or
gastrointestinal adverse events (see section 4.2).

Distribution

The apparent volume of distribution following oral administration of 240 mg Tecfidera varies between
60 L and 90 L. Human plasma protein binding of monomethyl fumarate generally ranges between
27% and 40%.

Biotransformation

In humans, dimethyl fumarate is extensively metabolised with less than 0.1% of the dose excreted as
unchanged dimethyl fumarate in urine. It is initially metabolised by esterases, which are ubiquitous in
the gastrointestinal tract, blood and tissues, before it reaches the systemic circulation. Further
metabolism occurs through the tricarboxylic acid cycle, with no involvement of the cytochrome P450
(CYP) system. A single 240 mg $^{14}$C-dimethyl fumarate dose study identified glucose as the
predominant metabolite in human plasma. Other circulating metabolites included fumaric acid, citric
acid and monomethyl fumarate. The downstream metabolism of fumaric acid occurs through the
tricarboxylic acid cycle, with exhalation of CO$_2$ serving as a primary route of elimination.

Elimination

Exhalation of CO$_2$ is the primary route of dimethyl fumarate elimination accounting for 60% of the
dose. Renal and faecal elimination are secondary routes of elimination, accounting for 15.5% and
0.9% of the dose respectively.

The terminal half-life of monomethyl fumarate is short (approximately 1 hour) and no circulating
monomethyl fumarate is present at 24 hours in the majority of individuals. Accumulation of parent
drug or monomethyl fumarate does not occur with multiple doses of dimethyl fumarate at the
therapeutic regimen.

Linearity

Dimethyl fumarate exposure increases in an approximately dose proportional manner with single and
multiple doses in the 120 mg to 360 mg dose range studied.

Pharmacokinetics in special patient groups

Based on the results of Analysis of Variance (ANOVA), body weight is the main covariate of
exposure (by $C_{\text{max}}$ and AUC) in relapsing remitting multiple sclerosis (RRMS) subjects, but did not
affect safety and efficacy measures evaluated in the clinical studies.

Gender and age did not have a clinically significant impact on the pharmacokinetics of dimethyl
fumarate. The pharmacokinetics in patients aged 65 and over has not been studied.

Paediatric population

The pharmacokinetics in patients below the age of 18 has not been studied.
**Renal impairment**

Since the renal pathway is a secondary route of elimination for dimethyl fumarate accounting for less than 16% of the dose administered, evaluation of pharmacokinetics in individuals with renal impairment was not conducted.

**Hepatic impairment**

As dimethyl fumarate and monomethyl fumarate are metabolised by esterases, without the involvement of the CYP450 system, evaluation of pharmacokinetics in individuals with hepatic impairment was not conducted.

### 5.3 Preclinical safety data

The adverse reactions described in the Toxicology and Reproduction toxicity sections below were not observed in clinical studies, but were seen in animals at exposure levels similar to clinical exposure levels.

**Mutagenesis**

Dimethyl fumarate and mono-methylfumarate were negative in a battery of *in vitro* assays (Ames, chromosomal aberration in mammalian cells). Dimethyl fumarate was negative in the *in vivo* micronucleus assay in the rat.

**Carcinogenesis**

Carcinogenicity studies of dimethyl fumarate were conducted for up to 2 years in mice and rats. Dimethyl fumarate was administered orally at doses of 25, 75, 200 and 400 mg/kg/day in mice, and at doses of 25, 50, 100, and 150 mg/kg/day in rats. In mice, the incidence of renal tubular carcinoma was increased at 75 mg/kg/day, at equivalent exposure (AUC) to the recommended human dose. In rats, the incidence of renal tubular carcinoma was increased at 100 mg/kg/day, approximately 3 times higher exposure than the recommended human dose. The relevance of these findings to human risk is unknown.

The incidence of squamous cell papilloma and carcinoma in the nonglandular stomach (forestomach) was increased at equivalent exposure to the recommended human dose in mice and below exposure to the recommended human dose in rats (based on AUC). The forestomach in rodents does not have a human counterpart.

**Toxicology**

Nonclinical studies in rodent, rabbits, and monkeys were conducted with a dimethyl fumarate suspension (dimethyl fumarate in 0.8% hydroxypropyl methylcellulose) administered by oral gavage. The chronic dog study was conducted with oral administration of the dimethyl fumarate capsule.

Kidney changes were observed after repeated oral administration of dimethyl fumarate in mice, rats, dogs, and monkeys. Renal tubule epithelial regeneration, suggestive of injury, was observed in all species. Renal tubular hyperplasia was observed in rats with life time dosing (2-year study). Cortical atrophy was observed in dogs and monkeys, and single cell necrosis and interstitial fibrosis were observed in monkeys that received daily oral doses of dimethyl fumarate for 12 months, at 6 times the recommended dose based on AUC. The relevance of these findings to humans is not known.

In the testes, degeneration of the seminiferous epithelium was seen in rats and dogs. The findings were observed at approximately the recommended dose in rats and 6 times the recommended dose in dogs (AUC basis). The relevance of these findings to humans is not known.

Findings in the forestomach of mice and rats consisted of squamous epithelial hyperplasia and
hyperkeratosis; inflammation; and squamous cell papilloma and carcinoma in studies of 3 months or longer in duration. The forestomach of mice and rats does not have a human counterpart.

**Reproduction toxicity**

Oral administration of dimethyl fumarate to male rats at 75, 250, and 375 mg/kg/day prior to and during mating had no effects on male fertility up to the highest dose tested (at least 2 times the recommended dose on an AUC basis). Oral administration of dimethyl fumarate to female rats at 25, 100, and 250 mg/kg/day prior to and during mating, and continuing to Day 7 of gestation, induced reduction in the number of estrous stages per 14 days and increased the number of animals with prolonged diestrus at the highest dose tested (11 times the recommended dose on an AUC basis). However, these changes did not affect fertility or the number of viable fetuses produced.

Dimethyl fumarate has been shown to cross the placental membrane into fetal blood in rats and rabbits, with ratios of fetal to maternal plasma concentrations of 0.48 to 0.64 and 0.1 respectively. No malformations were observed at any dose of dimethyl fumarate in rats or rabbits. Administration of dimethyl fumarate at oral doses of 25, 100, and 250 mg/kg/day to pregnant rats during the period of organogenesis resulted in maternal adverse effects at 4 times the recommended dose on an AUC basis, and low fetal weight and delayed ossification (metatarsals and hindlimb phalanges) at 11 times the recommended dose on an AUC basis. The lower fetal weight and delayed ossification were considered secondary to maternal toxicity (reduced body weight and food consumption).

Oral administration of dimethyl fumarate at 25, 75, and 150 mg/kg/day to pregnant rabbits during organogenesis had no effect on embryo-fetal development and resulted in reduced maternal body weight at 7 times the recommended dose and increased abortion at 16 times the recommended dose, on an AUC basis.

Oral administration of dimethyl fumarate at 25, 100, and 250 mg/kg/day to rats during pregnancy and lactation resulted in lower body weights in the F1 offspring, and delays in sexual maturation in F1 males at 11 times the recommended dose on an AUC basis. There were no effects on fertility in the F1 offspring. The lower offspring body weight was considered secondary to maternal toxicity.

### 6. PHARMACEUTICAL PARTICULARS

#### 6.1 List of excipients

**Enteric-coated microtablets**

- Microcrystalline cellulose
- Croscarmellose sodium
- Talc
- Silica, hydrophobic colloidal
- Magnesium stearate
- Triethyl citrate
- Methacrylic acid – methyl methacrylate copolymer (1:1)
- Methacrylic acid – ethyl acrylate copolymer (1:1) dispersion 30%
- Simeticone
- Sodium laurilsulfate
- Polysorbate 80

**Capsule shell**

- Gelatin
- Titanium dioxide (E171)
- Brilliant Blue FCF (E133)
- Yellow iron oxide (E172)
Capsule print (black ink)

Shellac
Potassium hydroxide
Black iron oxide (E172)

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

120 mg gastro-resistant hard capsules: 4 years

6.4 Special precautions for storage

Do not store above 30°C.
Keep the blisters in the outer carton in order to protect from light.

6.5 Nature and contents of container

120 mg capsules: 14 capsules in PVC/PE/PVDC-PVC aluminium blister packs.
Not all pack sizes may be marketed.

6.6 Special precautions for disposal

No special requirements.

7. MARKETING AUTHORISATION HOLDER

Biogen Idec Ltd
Innovation House
70 Norden Road
Maidenhead
Berkshire
SL6 4AY
United Kingdom

8. MARKETING AUTHORISATION NUMBER(S)

EU/1/13/837/001

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

Date of first authorisation: 30 January 2014

10. DATE OF REVISION OF THE TEXT

{MM/YYYY}

Detailed information on this medicinal product is available on the website of the European Medicines
1. NAME OF THE MEDICINAL PRODUCT
Tecfidera 240 mg gastro-resistant hard capsules

2. QUALITATIVE AND QUANTITATIVE COMPOSITION
Each capsule contains 240 mg dimethyl fumarate.
For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM
Gastro-resistant hard capsule
Green gastro-resistant hard capsule printed with ‘BG-12 240 mg’.

4. CLINICAL PARTICULARS
4.1 Therapeutic indications
Tecfidera is indicated for the treatment of adult patients with relapsing remitting multiple sclerosis (please refer to section 5.1 for important information on the populations for which efficacy has been established).

4.2 Posology and method of administration
Treatment should be initiated under supervision of a physician experienced in the treatment of the disease.

Posology
The starting dose is 120 mg twice a day. After 7 days, the dose is increased to the recommended dose of 240 mg twice a day.

Temporary dose reduction to 120 mg twice a day may reduce the occurrence of flushing and gastrointestinal adverse reactions. Within 1 month, the recommended dose of 240 mg twice a day should be resumed.

Tecfidera should be taken with food (see section 5.2). For those patients who may experience flushing or gastrointestinal adverse reactions, taking Tecfidera with food may improve tolerability (see sections 4.4, 4.5 and 4.8).

Older people
Clinical studies of Tecfidera had limited exposure to patients aged 55 years and above, and did not include sufficient numbers of patients aged 65 and over to determine whether they respond differently than younger patients (see section 5.2). Based on the mode of action of the active substance there are no theoretical reasons for any requirement for dose adjustments in the elderly.
Renal and hepatic impairment

Tecfidera has not been studied in patients with renal or hepatic impairment. Based on clinical pharmacology studies, no dose adjustments are needed (see section 5.2). Caution should be used when treating patients with severe renal or severe hepatic impairment (see section 4.4).

Paediatric population

The safety and efficacy of Tecfidera in children and adolescents aged 10 to 18 years have not been established. No data are available. There is no relevant use of Tecfidera in children aged less than 10 years in multiple sclerosis.

Method of administration

For oral use.

The capsule or its contents should not be crushed, divided, dissolved, sucked or chewed as the enteric-coating of the microtablets prevents irritant effects on the gut.

4.3 Contraindications

Hypersensitivity to the active substance or to any of the excipients listed in section 6.1.

4.4 Special warnings and precautions for use

Blood/laboratory tests

Tecfidera may decrease lymphocyte counts (see section 4.8). Tecfidera has not been studied in patients with pre-existing low lymphocyte counts and caution should be exercised when treating these patients. Prior to initiating treatment with Tecfidera, a recent complete blood count (i.e. within 6 months) should be available. Assessments of complete blood counts are also recommended after 6 months of treatment and every 6 to 12 months thereafter and as clinically indicated.

Changes in renal and hepatic laboratory tests have been seen in clinical trials in subjects treated with Tecfidera (see section 4.8). The clinical implications of these changes are unknown. Assessments of renal function (e.g. creatinine, blood urea nitrogen and urinalysis) and hepatic function (e.g. ALT and AST) are recommended prior to treatment initiation, after 3 and 6 months of treatment, every 6 to 12 months thereafter and as clinically indicated.

Severe renal and hepatic impairment

Tecfidera has not been studied in patients with severe renal or severe hepatic impairment and caution should, therefore, be used in these patients (see section 4.2).

Severe active gastrointestinal disease

Tecfidera has not been studied in patients with severe active gastrointestinal disease and caution should, therefore, be used in these patients.

Flushing

In clinical trials, 34% of Tecfidera treated patients experienced flushing. In the majority of patients who experienced flushing, it was mild or moderate in severity.

In clinical trials, 3 patients out of a total of 2,560 patients treated with Tecfidera experienced serious flushing symptoms that were probable hypersensitivity or anaphylactoid reactions. These events were not life-threatening, but led to hospitalisation. Prescribers and patients should be alert to this
possibility in the event of severe flushing reactions (see sections 4.2, 4.5 and 4.8).

Infections

In phase III placebo-controlled studies, the incidence of infections (60% vs 58%) and serious infections (2% vs 2%) was similar in patients treated with Tecfidera or placebo, respectively. There was no increased incidence of serious infections observed in patients with lymphocyte counts <0.8x10^9/L or <0.5x10^9/L. During treatment with Tecfidera in the MS placebo controlled trials, mean lymphocyte counts decreased by approximately 30% from baseline at one year and then plateaued (see section 4.8). Mean lymphocyte counts remained within normal limits. If a patient develops a serious infection, suspending treatment with Tecfidera should be considered and the benefits and risks should be reassessed prior to re-initiation of therapy. Patients receiving Tecfidera should be instructed to report symptoms of infections to a physician. Patients with serious infections should not start treatment with Tecfidera until the infection(s) is resolved.

4.5 Interaction with other medicinal products and other forms of interaction

Tecfidera has not been studied in combination with anti-neoplastic or immunosuppressive therapies and caution should, therefore, be used during concomitant administration. In multiple sclerosis clinical studies, the concomitant treatment of relapses with a short course of intravenous corticosteroids was not associated with a clinically relevant increase of infection.

Vaccination during treatment with Tecfidera has not been studied. It is not known whether treatment with Tecfidera might reduce the effectiveness of some vaccines. Live vaccines might carry an increased risk of clinical infection and should not be given to patients treated with Tecfidera unless, in exceptional cases, this potential risk is considered to be outweighed by the risk to the individual of not vaccinating.

During treatment with Tecfidera, simultaneous use of other fumaric acid derivatives (topical or systemic) should be avoided.

In humans, dimethyl fumarate is extensively metabolised by esterases before it reaches the systemic circulation and further metabolism occurs through the tricarboxylic acid cycle, with no involvement of the cytochrome P450 (CYP) system. Potential drug interaction risks were not identified from in vitro CYP-inhibition and induction studies, a p-glycoprotein study, or studies of the protein binding of dimethyl fumarate and monomethyl fumarate (a primary metabolite of dimethyl fumarate).

Commonly used medicinal products in patients with multiple sclerosis, intramuscular interferon beta-1a and glatiramer acetate, were clinically tested for potential interactions with dimethyl fumarate and did not alter the pharmacokinetic profile of dimethyl fumarate.

Administration of 325 mg (or equivalent) non-enteric coated acetylsalicylic acid, 30 minutes prior to Tecfidera, over 4 days of dosing, did not alter the pharmacokinetic profile of Tecfidera and reduced the occurrence and severity of flushing in a healthy volunteer study. However, long term use of acetylsalicylic acid is not recommended for the management of flushing. Potential risks associated with acetylsalicylic acid therapy should be considered prior to co-administration with Tecfidera. (see sections 4.2, 4.4 and 4.8).

Concurrent therapy with nephrotoxic medicinal products (such as aminoglycosides, diuretics, NSAIDs or lithium) may increase the potential of renal adverse reactions (e.g. proteinuria) in patients taking Tecfidera (see section 4.8).

Consumption of moderate amounts of alcohol did not alter exposure to Tecfidera and was not associated with an increase in adverse reactions. Consumption of large quantities of undiluted strong alcoholic drinks (more than 30% alcohol by volume) may lead to increased dissolution rates of Tecfidera and, therefore, may increase the frequency of gastrointestinal adverse reactions.
In vitro CYP induction studies did not demonstrate an interaction between Tecfidera and oral contraceptives. In vivo interaction studies have not been performed with oral contraceptives. Even though an interaction is not expected, non-hormonal contraceptive measures should be considered with Tecfidera (see section 4.6).

Paediatric population

Interaction studies have only been performed in adults.

4.6 Fertility, pregnancy and lactation

Pregnancy

There are no or limited amount of data from the use of dimethyl fumarate in pregnant women. Animal studies have shown reproductive toxicity (see section 5.3). Tecfidera is not recommended during pregnancy and in women of childbearing potential not using appropriate contraception (see section 4.5). Tecfidera should be used during pregnancy only if clearly needed and if the potential benefit justifies the potential risk to the foetus.

Breast-feeding

It is unknown whether dimethyl fumarate or its metabolites are excreted in human milk. A risk to the newborns/infants cannot be excluded. A decision must be made whether to discontinue breast-feeding or to discontinue Tecfidera therapy. The benefit of breast-feeding for the child and the benefit of therapy for the woman should be taken into account.

Fertility

There are no data on the effects of Tecfidera on human fertility. Data from preclinical studies do not suggest that dimethyl fumarate would be associated with an increased risk of reduced fertility (see section 5.3).

4.7 Effects on ability to drive and use machines

No studies on the ability to drive and use machines have been conducted.

4.8 Undesirable effects

Summary of the safety profile

The most common adverse reactions (incidence ≥10%) for patients treated with Tecfidera were flushing and gastrointestinal events (i.e. diarrhoea, nausea, abdominal pain, abdominal pain upper). Flushing and gastrointestinal events tend to begin early in the course of treatment (primarily during the first month) and in patients who experience flushing and gastrointestinal events, these events may continue to occur intermittently throughout treatment with Tecfidera. The most commonly reported adverse reactions leading to discontinuation (incidence >1%) in patients treated with Tecfidera were flushing (3%) and gastrointestinal events (4%).

In placebo-controlled and uncontrolled clinical studies, a total of 2,468 patients have received Tecfidera and been followed for periods up to 4 years with an overall exposure equivalent to 3,588 person-years. Approximately 1,056 patients have received more than 2 years of treatment with Tecfidera. The experience in uncontrolled clinical trials is consistent with the experience in the placebo-controlled clinical trials.
Tabulated summary of adverse reactions

Adverse reactions, which were more frequently reported in Tecfidera versus placebo-treated patients, are presented in the table below. These data were derived from 2 pivotal Phase 3 placebo-controlled, double-blind clinical trials with a total of 1,529 patients treated with Tecfidera and for up to 24 months with an overall exposure of 2,371 person-years (see section 5.1). The frequencies described in the table below are based on 769 patients treated with Tecfidera 240 mg twice a day and 771 patients treated with placebo.

The adverse reactions are presented as MedDRA preferred terms under the MedDRA System Organ Class. The incidence of the adverse reactions below are expressed according to the following categories:
- Very common (≥1/10)
- Common (≥1/100 to <1/10)
- Uncommon (≥1/1,000 to <1/100)
- Rare (≥1/10,000 to <1/1,000)
- Very rare (<1/10,000)
- Not known (cannot be estimated from the available data)

<table>
<thead>
<tr>
<th>MedDRA System Organ Class</th>
<th>Adverse reaction</th>
<th>Frequency category</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infections and infestations</td>
<td>Gastroenteritis</td>
<td>Common</td>
</tr>
<tr>
<td>Blood and lymphatic system disorders</td>
<td>Lymphopenia</td>
<td>Common</td>
</tr>
<tr>
<td></td>
<td>Leucopenia</td>
<td>Common</td>
</tr>
<tr>
<td>Immune system disorders</td>
<td>Hypersensitivity</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Nervous system disorders</td>
<td>Burning sensation</td>
<td>Common</td>
</tr>
<tr>
<td>Vascular disorders</td>
<td>Flushing</td>
<td>Very common</td>
</tr>
<tr>
<td></td>
<td>Hot flush</td>
<td>Common</td>
</tr>
<tr>
<td>Gastrointestinal disorders</td>
<td>Diarrhoea</td>
<td>Very common</td>
</tr>
<tr>
<td></td>
<td>Nausea</td>
<td>Very common</td>
</tr>
<tr>
<td></td>
<td>Abdominal pain upper</td>
<td>Very common</td>
</tr>
<tr>
<td></td>
<td>Abdominal pain</td>
<td>Very common</td>
</tr>
<tr>
<td></td>
<td>Vomiting</td>
<td>Common</td>
</tr>
<tr>
<td></td>
<td>Dyspepsia</td>
<td>Common</td>
</tr>
<tr>
<td></td>
<td>Gastritis</td>
<td>Common</td>
</tr>
<tr>
<td></td>
<td>Gastrointestinal disorder</td>
<td>Common</td>
</tr>
<tr>
<td>Skin and subcutaneous tissue disorders</td>
<td>Pruritus</td>
<td>Common</td>
</tr>
<tr>
<td></td>
<td>Rash</td>
<td>Common</td>
</tr>
<tr>
<td></td>
<td>Erythema</td>
<td>Common</td>
</tr>
<tr>
<td>Renal and urinary disorders</td>
<td>Proteinuria</td>
<td>Common</td>
</tr>
<tr>
<td>General disorders and administration site conditions</td>
<td>Feeling hot</td>
<td>Common</td>
</tr>
<tr>
<td>Investigations</td>
<td>Ketones measured in urine</td>
<td>Very common</td>
</tr>
<tr>
<td></td>
<td>Albumin urine present</td>
<td>Common</td>
</tr>
<tr>
<td></td>
<td>Aspartate aminotransferase increased</td>
<td>Common</td>
</tr>
<tr>
<td></td>
<td>Alanine aminotransferase increased</td>
<td>Common</td>
</tr>
<tr>
<td></td>
<td>White blood cell count decreased</td>
<td>Common</td>
</tr>
</tbody>
</table>

Description of selected adverse reactions

Flushing

In the placebo-controlled studies, the incidence of flushing (34% versus 4%) and hot flush (7% versus 2%) was increased in patients treated with Tecfidera compared to placebo, respectively. Flushing is usually described as flushing or hot flush, but can include other events (e.g. warmth, redness, itching, and burning sensation). Flushing events tend to begin early in the course of treatment (primarily during the first month) and in patients who experience flushing, these events may continue
to occur intermittently throughout treatment with Tecfidera. In patients with flushing, the majority had flushing events that were mild or moderate in severity. Overall, 3% of patients treated with Tecfidera discontinued due to flushing. The incidence of serious flushing, which may be characterised by generalised erythema, rash and/or pruritus, was seen in less than 1% of patients treated with Tecfidera (see sections 4.2, 4.4 and 4.5).

**Gastrointestinal**

The incidence of gastrointestinal events (e.g. diarrhoea [14% versus 10%], nausea [12% versus 9%], upper abdominal pain [10% versus 6%], abdominal pain [9% versus 4%], vomiting [8% versus 5%] and dyspepsia [5% versus 3%]) was increased in patients treated with Tecfidera compared to placebo, respectively. Gastrointestinal events tend to begin early in the course of treatment (primarily during the first month) and in patients who experience gastrointestinal events, these events may continue to occur intermittently throughout treatment with Tecfidera. In the majority of patients who experienced gastrointestinal events, it was mild or moderate in severity. Four per cent (4%) of patients treated with Tecfidera discontinued due to gastrointestinal events. The incidence of serious gastrointestinal events, including gastroenteritis and gastritis, was seen in 1% of patients treated with Tecfidera (see section 4.2).

**Hepatic transaminases**

In placebo-controlled studies, elevations of hepatic transaminases were observed. The majority of patients with elevations had hepatic transaminases that were <3 times the upper limit of normal (ULN). The increased incidence of elevations of hepatic transaminases in patients treated with Tecfidera relative to placebo was primarily seen during the first 6 months of treatment. Elevations of alanine aminotransferase and aspartate aminotransferase ≥3 times ULN, respectively, were seen in 5% and 2% of patients treated with placebo and 6% and 2% of patients treated with Tecfidera. There were no elevations in transaminases ≥3 times ULN with concomitant elevations in total bilirubin >2 times ULN. Discontinuations due to elevated hepatic transaminases were <1% and similar in patients treated with Tecfidera or placebo.

**Renal**

In placebo-controlled studies, the incidence of proteinuria was higher in patients treated with Tecfidera (9%) compared to placebo (7%). The overall incidence of renal and urinary adverse events was similar for Tecfidera and placebo-treated patients. There were no reports of serious renal failure. On urinalysis, the percentage of patients with protein values of 1+ or greater was similar for Tecfidera (43%) and placebo-treated patients (40%). Typically, laboratory observations of proteinuria were not progressive. Compared to patients treated with placebo, estimated glomerular filtration rate (eGFR) was observed to increase in patients treated with Tecfidera, including those patients with 2 consecutive occurrences of proteinuria (≥1+).

**Haematological**

In the placebo-controlled studies, most patients (>98%) had normal lymphocyte values prior to initiating treatment. Upon treatment with Tecfidera, mean lymphocyte counts decreased over the first year with a subsequent plateau. On average, lymphocyte counts decreased by approximately 30% of baseline value. Mean and median lymphocyte counts remained within normal limits. Lymphocyte counts <0.5x10⁹/l were observed in <1% of patients treated with placebo and 6% of patients treated with Tecfidera. A lymphocyte count <0.2x10⁹/l was observed in 1 patient treated with Tecfidera and in no patients treated with placebo. The incidence of infections (58% versus 60%) and serious infections (2% versus 2%) was similar in patients treated with placebo or Tecfidera. An increased incidence of infections and serious infections was not observed in patients with lymphocyte counts <0.8x10⁹/l or <0.5x10⁹/l. A transient increase in mean eosinophil counts was seen during the first 2 months of therapy.
Laboratory abnormalities

In the placebo-controlled studies, measurement of urinary ketones (1+ or greater) was higher in patients treated with Tecfidera (45%) compared to placebo (10%). No untoward clinical consequences were observed in clinical trials.

Levels of 1,25-dihydroxyvitamin D decreased in Tecfidera treated patients relative to placebo (median percentage decrease from baseline at 2 years of 25% versus 15%, respectively) and levels of parathyroid hormone (PTH) increased in Tecfidera treated patients relative to placebo (median percentage increase from baseline at 2 years of 29% versus 15%, respectively). Mean values for both parameters remained within normal range.

Reporting of suspected adverse reactions

Reporting suspected adverse reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/risk balance of the medicinal product. Healthcare professionals are asked to report any suspected adverse reactions via the national reporting system listed in Appendix V.

4.9 Overdose

No cases of overdose have been reported.

5. Pharmacological properties

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Other nervous system drugs, ATC code: N07XX09

Mechanism of action

The mechanism by which dimethyl fumarate exerts therapeutic effects in multiple sclerosis is not fully understood. Preclinical studies indicate that dimethyl fumarate pharmacodynamic responses appear to be primarily mediated through activation of the Nuclear factor (erythroid-derived 2)-like 2 (Nrf2) transcriptional pathway. Dimethyl fumarate has been shown to up regulate Nrf2-dependent antioxidant genes in patients (e.g. NAD(P)H dehydrogenase, quinone 1; [NQO1]).

Pharmacodynamic effects

Effects on the immune system

In preclinical and clinical studies, Tecfidera demonstrated anti-inflammatory and immunomodulatory properties. Dimethyl fumarate and monomethyl fumarate, the primary metabolite of dimethyl fumarate, significantly reduced immune cell activation and subsequent release of pro-inflammatory cytokines in response to inflammatory stimuli in preclinical models. In clinical studies with psoriasis patients, dimethyl fumarate affected lymphocyte phenotypes through a down-regulation of pro-inflammatory cytokine profiles (T_{H1}, T_{H17}), and biased towards anti-inflammatory production (T_{H2}). Dimethyl fumarate demonstrated therapeutic activity in multiple models of inflammatory and neuroinflammatory injury. In Phase 3 studies, upon treatment with Tecfidera mean lymphocyte counts decreased on average by approximately 30% of their baseline value over the first year with a subsequent plateau.

Effect on cardiovascular system

Single doses of 240 mg or 360 mg Tecfidera did not have any effect on the QTc interval when compared to placebo in a QTc study.
Clinical efficacy and safety

Two, 2-year, randomised, double-blind, placebo controlled studies [Study 1 (DEFINE) with 1234 subjects and Study 2 (CONFIRM) with 1417 subjects] of subjects with relapsing-remitting multiple sclerosis (RRMS) were performed. Subjects with progressive forms of MS were not included in these studies. Efficacy (see table below) and safety were demonstrated in subjects with Expanded Disability Status Scale (EDSS) scores ranging from 0 to 5 inclusive, who had experienced at least 1 relapse during the year prior to randomisation, or, within 6 weeks of randomisation had a brain Magnetic Resonance Imaging (MRI) demonstrating at least one gadolinium-enhancing (Gd+) lesion. Study 2 contained a rater-blinded (i.e. study physician/ investigator assessing the response to study treatment was blinded) reference comparator of glatiramer acetate.

In Study 1, patients had the following median baseline characteristics: age 39 years, disease duration 7.0 years, EDSS score 2.0. In addition, 16% of patients had an EDSS score >3.5, 28% had ≥2 relapses in the prior year and 42% had previously received other approved MS treatments. In the MRI cohort 36% of patients entering the study had Gd+ lesions at baseline (mean number of Gd+ lesions 1.4).

In Study 2, patients had the following median baseline characteristics: age 37 years, disease duration 6.0 years, EDSS score 2.5. In addition, 17% of patients had an EDSS score >3.5, 32% had ≥2 relapses in the prior year and 30% had previously received other approved MS treatments. In the MRI cohort 45% of patients entering the study had Gd+ lesions at baseline (mean number of Gd+ lesions 2.4).

Compared to placebo, subjects treated with Tecfidera had a clinically meaningful and statistically significant reduction on: the primary endpoint in Study 1, proportion of subjects relapsed at 2 years; and the primary endpoint in Study 2, annualised relapse rate at 2 years.

The annualised relapse rate for glatiramer acetate and placebo was 0.286 and 0.401 respectively in Study 2, corresponding to a reduction of 29% (p=0.013), which is consistent with approved prescribing information.

<table>
<thead>
<tr>
<th>Clinical Endpoints</th>
<th>DEFINE</th>
<th>CONFIRM</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Placebo</td>
<td>Tecfidera 240 mg twice a day</td>
</tr>
<tr>
<td>No. subjects</td>
<td>408</td>
<td>410</td>
</tr>
<tr>
<td>Annualised relapse rate</td>
<td>0.364</td>
<td>0.172***</td>
</tr>
<tr>
<td>Rate ratio (95% CI)</td>
<td>0.47 (0.37, 0.61)</td>
<td>0.56 (0.42, 0.74)</td>
</tr>
<tr>
<td>Proportion relapsed</td>
<td>0.461</td>
<td>0.270***</td>
</tr>
<tr>
<td>Hazard ratio (95% CI)</td>
<td>0.51 (0.40, 0.66)</td>
<td>0.66 (0.51, 0.86)</td>
</tr>
<tr>
<td>Proportion with 12-week confirmed disability progression</td>
<td>0.271</td>
<td>0.164**</td>
</tr>
<tr>
<td>Hazard ratio (95% CI)</td>
<td>0.62 (0.44, 0.87)</td>
<td>0.79 (0.52, 1.19)</td>
</tr>
<tr>
<td>Proportion with 24 week confirmed disability progression</td>
<td>0.169</td>
<td>0.128#</td>
</tr>
<tr>
<td>Hazard ratio (95% CI)</td>
<td>0.77 (0.52, 1.14)</td>
<td>0.62 (0.37, 1.03)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>MRI Endpoints</th>
<th>DEFINE</th>
<th>CONFIRM</th>
<th>Glatiramer acetate</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. subjects</td>
<td>165</td>
<td>152</td>
<td>144 147 161</td>
</tr>
<tr>
<td>DEFINE</td>
<td>CONFIRM</td>
<td></td>
<td></td>
</tr>
<tr>
<td>--------</td>
<td>---------</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Placebo</td>
<td>Tecfidera 240 mg twice a day</td>
</tr>
<tr>
<td>Mean (median) number of new or newly enlarging T2 lesions over 2 years</td>
<td>16.5 (7.0)***</td>
<td>3.2 (1.0)***</td>
<td>19.9 (11.0)***</td>
</tr>
<tr>
<td>Lesion mean ratio (95% CI)</td>
<td>0.15 (0.10, 0.23)</td>
<td>0.29 (0.21, 0.41)</td>
<td>0.46 (0.33, 0.63)</td>
</tr>
<tr>
<td>Mean (median) number of Gd lesions at 2 years</td>
<td>1.8 (0)***</td>
<td>0.1 (0)***</td>
<td>2.0 (0)***</td>
</tr>
<tr>
<td>Odds ratio (95% CI)</td>
<td>0.10 (0.05, 0.22)</td>
<td>0.26 (0.15, 0.46)</td>
<td>0.39 (0.24, 0.65)</td>
</tr>
<tr>
<td>Mean (median) number of new T1 hypointense lesions over 2 years</td>
<td>5.7 (2.0)***</td>
<td>2.0 (1.0)***</td>
<td>8.1 (4.0)***</td>
</tr>
<tr>
<td>Lesion mean ratio (95% CI)</td>
<td>0.28 (0.20, 0.39)</td>
<td>0.43 (0.30, 0.61)</td>
<td>0.59 (0.42, 0.82)</td>
</tr>
</tbody>
</table>

<sup>a</sup>All analyses of clinical endpoints were intent-to-treat; <sup>b</sup>MRI analysis used MRI cohort

*P-value < 0.05; **P-value < 0.01; ***P-value < 0.0001; #not statistically significant

Efficacy in patients with high disease activity:
Consistent treatment effect on relapses in a subgroup of patients with high disease activity was observed, whilst the effect on time to 3-month sustained disability progression was not clearly established. Due to the design of the studies, high disease activity was defined as follows:
- Patients with 2 or more relapses in one year, and with one or more Gd-enhancing lesions on brain MRI (n=42 in DEFINE; n=51 in CONFIRM) or,
- Patients who have failed to respond to a full and adequate course (at least one year of treatment) of beta-interferon, having had at least 1 relapse in the previous year while on therapy, and at least 9 T2-hyperintense lesions in cranial MRI or at least 1 Gd-enhancing lesion, or patients having an unchanged or increased relapse rate in the prior year as compared to the previous 2 years (n=177 in DEFINE; n=141 in CONFIRM).

Paediatric population

The European Medicines Agency has deferred the obligation to submit the results of studies with Tecfidera in one or more subsets of the paediatric population in multiple sclerosis (see section 4.2 for information on paediatric use).

### 5.2 Pharmacokinetic properties

Orally administered Tecfidera (dimethyl fumarate) undergoes rapid presystemic hydrolysis by esterases and is converted to its primary metabolite, monomethyl fumarate, which is also active. Dimethyl fumarate is not quantifiable in plasma following oral administration of Tecfidera. Therefore, all pharmacokinetic analyses related to dimethyl fumarate were performed with plasma monomethyl fumarate concentrations. Pharmacokinetic data were obtained in subjects with multiple sclerosis and healthy volunteers.

#### Absorption

The $T_{max}$ of monomethyl fumarate is 2 to 2.5 hours. As Tecfidera gastro-resistant hard capsules contain microtablets, which are protected by an enteric coating, absorption does not commence until they leave the stomach (generally less than 1 hour). Following 240 mg twice a day administered with food, the median peak ($C_{max}$) was 1.72 mg/l and overall (AUC) exposure was 8.02 h.mg/l in subjects with multiple sclerosis. Overall, $C_{max}$ and AUC increased approximately dose-proportionally in the dose range studied (120 mg to 360 mg). In subjects with multiple sclerosis, two 240 mg doses were administered 4 hours apart as part of a three times a day dosing regimen. This resulted in a minimal
accumulation of exposure yielding an increase in the median Cmax of 12% compared to the twice
daily dosing (1.72 mg/l for twice daily compared to 1.93 mg/l for three times daily) with no safety
implications.

Food does not have a clinically significant effect on exposure of dimethyl fumarate. However,
Tecfidera should be taken with food due to improved tolerability with respect to flushing or
gastrointestinal adverse events (see section 4.2).

Distribution

The apparent volume of distribution following oral administration of 240 mg Tecfidera varies between
60 L and 90 L. Human plasma protein binding of monomethyl fumarate generally ranges between
27% and 40%.

Biotransformation

In humans, dimethyl fumarate is extensively metabolised with less than 0.1% of the dose excreted as
unchanged dimethyl fumarate in urine. It is initially metabolised by esterases, which are ubiquitous in
the gastrointestinal tract, blood and tissues, before it reaches the systemic circulation. Further
metabolism occurs through the tricarboxylic acid cycle, with no involvement of the cytochrome P450
(CYP) system. A single 240 mg 14C-dimethyl fumarate dose study identified glucose as the
predominant metabolite in human plasma. Other circulating metabolites included fumaric acid, citric
acid and monomethyl fumarate. The downstream metabolism of fumaric acid occurs through the
tricarboxylic acid cycle, with exhalation of CO₂ serving as a primary route of elimination.

Elimination

Exhalation of CO₂ is the primary route of dimethyl fumarate elimination accounting for 60% of the
dose. Renal and faecal elimination are secondary routes of elimination, accounting for 15.5% and
0.9% of the dose respectively.

The terminal half-life of monomethyl fumarate is short (approximately 1 hour) and no circulating
monomethyl fumarate is present at 24 hours in the majority of individuals. Accumulation of parent
drug or monomethyl fumarate does not occur with multiple doses of dimethyl fumarate at the
therapeutic regimen.

Linearity

Dimethyl fumarate exposure increases in an approximately dose proportional manner with single and
multiple doses in the 120 mg to 360 mg dose range studied.

Pharmacokinetics in special patient groups

Based on the results of Analysis of Variance (ANOVA), body weight is the main covariate of
exposure (by Cmax and AUC) in relapsing remitting multiple sclerosis (RRMS) subjects, but did not
affect safety and efficacy measures evaluated in the clinical studies.

Gender and age did not have a clinically significant impact on the pharmacokinetics of dimethyl
fumarate. The pharmacokinetics in patients aged 65 and over has not been studied.

Paediatric population

The pharmacokinetics in patients below the age of 18 has not been studied.
Renal impairment

Since the renal pathway is a secondary route of elimination for dimethyl fumarate accounting for less than 16% of the dose administered, evaluation of pharmacokinetics in individuals with renal impairment was not conducted.

Hepatic impairment

As dimethyl fumarate and monomethyl fumarate are metabolised by esterases, without the involvement of the CYP450 system, evaluation of pharmacokinetics in individuals with hepatic impairment was not conducted.

5.3 Preclinical safety data

The adverse reactions described in the Toxicology and Reproduction toxicity sections below were not observed in clinical studies, but were seen in animals at exposure levels similar to clinical exposure levels.

Mutagenesis

Dimethyl fumarate and mono-methylfumarate were negative in a battery of in vitro assays (Ames, chromosomal aberration in mammalian cells). Dimethyl fumarate was negative in the in vivo micronucleus assay in the rat.

Carcinogenesis

Carcinogenicity studies of dimethyl fumarate were conducted for up to 2 years in mice and rats. Dimethyl fumarate was administered orally at doses of 25, 75, 200 and 400 mg/kg/day in mice, and at doses of 25, 50, 100, and 150 mg/kg/day in rats. In mice, the incidence of renal tubular carcinoma was increased at 75 mg/kg/day, at equivalent exposure (AUC) to the recommended human dose. In rats, the incidence of renal tubular carcinoma was increased at 100 mg/kg/day, approximately 3 times higher exposure than the recommended human dose. The relevance of these findings to human risk is unknown.

The incidence of squamous cell papilloma and carcinoma in the nonglandular stomach (forestomach) was increased at equivalent exposure to the recommended human dose in mice and below exposure to the recommended human dose in rats (based on AUC). The forestomach in rodents does not have a human counterpart.

Toxicology

Nonclinical studies in rodent, rabbits, and monkeys were conducted with a dimethyl fumarate suspension (dimethyl fumarate in 0.8% hydroxypropyl methylcellulose) administered by oral gavage. The chronic dog study was conducted with oral administration of the dimethyl fumarate capsule.

Kidney changes were observed after repeated oral administration of dimethyl fumarate in mice, rats, dogs, and monkeys. Renal tubule epithelial regeneration, suggestive of injury, was observed in all species. Renal tubular hyperplasia was observed in rats with life time dosing (2-year study). Cortical atrophy was observed in dogs and monkeys, and single cell necrosis and interstitial fibrosis were observed in monkeys that received daily oral doses of dimethyl fumarate for 12 months, at 6 times the recommended dose based on AUC. The relevance of these findings to humans is not known.

In the testes, degeneration of the seminiferous epithelium was seen in rats and dogs. The findings were observed at approximately the recommended dose in rats and 6 times the recommended dose in dogs (AUC basis). The relevance of these findings to humans is not known.

Findings in the forestomach of mice and rats consisted of squamous epithelial hyperplasia and
hyperkeratosis; inflammation; and squamous cell papilloma and carcinoma in studies of 3 months or longer in duration. The forestomach of mice and rats does not have a human counterpart.

**Reproduction toxicity**

Oral administration of dimethyl fumarate to male rats at 75, 250, and 375 mg/kg/day prior to and during mating had no effects on male fertility up to the highest dose tested (at least 2 times the recommended dose on an AUC basis). Oral administration of dimethyl fumarate to female rats at 25, 100, and 250 mg/kg/day prior to and during mating, and continuing to Day 7 of gestation, induced reduction in the number of estrous stages per 14 days and increased the number of animals with prolonged diestrus at the highest dose tested (11 times the recommended dose on an AUC basis). However, these changes did not affect fertility or the number of viable fetuses produced.

Dimethyl fumarate has been shown to cross the placental membrane into fetal blood in rats and rabbits, with ratios of fetal to maternal plasma concentrations of 0.48 to 0.64 and 0.1 respectively. No malformations were observed at any dose of dimethyl fumarate in rats or rabbits. Administration of dimethyl fumarate at oral doses of 25, 100, and 250 mg/kg/day to pregnant rats during the period of organogenesis resulted in maternal adverse effects at 4 times the recommended dose on an AUC basis, and low fetal weight and delayed ossification (metatarsals and hindlimb phalanges) at 11 times the recommended dose on an AUC basis. The lower fetal weight and delayed ossification were considered secondary to maternal toxicity (reduced body weight and food consumption).

Oral administration of dimethyl fumarate at 25, 75, and 150 mg/kg/day to pregnant rabbits during organogenesis had no effect on embryo-fetal development and resulted in reduced maternal body weight at 7 times the recommended dose and increased abortion at 16 times the recommended dose, on an AUC basis.

Oral administration of dimethyl fumarate at 25, 100, and 250 mg/kg/day to rats during pregnancy and lactation resulted in lower body weights in the F1 offspring, and delays in sexual maturation in F1 males at 11 times the recommended dose on an AUC basis. There were no effects on fertility in the F1 offspring. The lower offspring body weight was considered secondary to maternal toxicity.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

**Enteric-coated microtablets**

- Microcrystalline cellulose
- Croscarmellose sodium
- Talc
- Silica, hydrophobic colloidal
- Magnesium stearate
- Triethyl citrate
- Methacrylic acid – methyl methacrylate copolymer (1:1)
- Methacrylic acid – ethyl acrylate copolymer (1:1) dispersion 30%
- Simeticone
- Sodium laurilsulfate
- Polysorbate 80

**Capsule shell**

- Gelatin
- Titanium dioxide (E171)
- Brilliant Blue FCF (E133)
- Yellow iron oxide (E172)
Capsule print (black ink)

Shellac
Potassium hydroxide
Black iron oxide (E172)

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

240 mg gastro-resistant hard capsules: 3 years

6.4 Special precautions for storage

Do not store above 30°C.
Keep the blisters in the outer carton in order to protect from light.

6.5 Nature and contents of container

240 mg capsules: 56 or 168 capsules in PVC/PE/PVDC-PVC aluminium blister packs.
Not all pack sizes may be marketed.

6.6 Special precautions for disposal

No special requirements.

7. MARKETING AUTHORISATION HOLDER

Biogen Idec Ltd
Innovation House
70 Norden Road
Maidenhead
Berkshire
SL6 4AY
United Kingdom

8. MARKETING AUTHORISATION NUMBER(S)

EU/1/13/837/002
EU/1/13/837/003

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

Date of first authorisation: 30 January 2014
10. DATE OF REVISION OF THE TEXT

{MM/YYYY}

Detailed information on this medicinal product is available on the website of the European Medicines Agency http://www.ema.europa.eu.
ANNEX II

A. MANUFACTURER RESPONSIBLE FOR BATCH RELEASE

B. CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE

C. OTHER CONDITIONS AND REQUIREMENTS OF THE MARKETING AUTHORISATION

D. CONDITIONS OR RESTRICTIONS WITH REGARD TO THE SAFE AND EFFECTIVE USE OF THE MEDICINAL PRODUCT
A. MANUFACTURER RESPONSIBLE FOR BATCH RELEASE

Name and address of the manufacturer responsible for batch release

Biogen Idec (Denmark) Manufacturing ApS
Biogen Idec Allé 1
DK-3400 Hillerod
Denmark

B. CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE

Medicinal product subject to restricted medical prescription (see Annex I: Summary of Product Characteristics, section 4.2).

C. OTHER CONDITIONS AND REQUIREMENTS OF THE MARKETING AUTHORISATION

- Periodic safety update reports

The marketing authorisation holder shall submit the first periodic safety update report for this product within 6 months following authorisation. Subsequently, the marketing authorisation holder shall submit periodic safety update reports for this product in accordance with the requirements set out in the list of Union reference dates (EURD list) provided for under Article 107c(7) of Directive 2001/83/EC and published on the European medicines web-portal.

D. CONDITIONS OR RESTRICTIONS WITH REGARD TO THE SAFE AND EFFECTIVE USE OF THE MEDICINAL PRODUCT

- Risk Management Plan (RMP)

The MAH shall perform the required pharmacovigilance activities and interventions detailed in the agreed RMP presented in Module 1.8.2 of the Marketing Authorisation and any agreed subsequent updates of the RMP.

An updated RMP should be submitted:

- At the request of the European Medicines Agency;
- Whenever the risk management system is modified, especially as the result of new information being received that may lead to a significant change to the benefit/risk profile or as the result of an important (pharmacovigilance or risk minimisation) milestone being reached.

If the dates for submission of a PSUR and the update of a RMP coincide, they can be submitted at the same time.
ANNEX III

LABELLING AND PACKAGE LEAFLET
A. LABELLING
**PARTICULARS TO APPEAR ON THE OUTER PACKAGING**

**OUTER CARTON**

1. **NAME OF THE MEDICINAL PRODUCT**

   Tecfidera 120 mg Gastro-resistant hard capsules  
   Dimethyl fumarate

2. **STATEMENT OF ACTIVE SUBSTANCE(S)**

   Each capsule contains 120 mg dimethyl fumarate

3. **LIST OF EXCIPIENTS**

4. **PHARMACEUTICAL FORM AND CONTENTS**

   14 gastro-resistant hard capsules

5. **METHOD AND ROUTE(S) OF ADMINISTRATION**

   Read the package leaflet before use  
   Oral use

6. **SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN**

   Keep out of the sight and reach of children

7. **OTHER SPECIAL WARNING(S), IF NECESSARY**

8. **EXPIRY DATE**

   EXP

9. **SPECIAL STORAGE CONDITIONS**

   Do not store above 30°C  
   Store in original package in order to protect from light

10. **SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE**
11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER

Biogen Idec Ltd,
Innovation House, 70 Norden Road,
Maidenhead, Berkshire, SL6 4AY
United Kingdom

12. MARKETING AUTHORISATION NUMBER(S)

EU/1/13/837/001

13. BATCH NUMBER

Lot

14. GENERAL CLASSIFICATION FOR SUPPLY

Medicinal product subject to medical prescription

15. INSTRUCTIONS ON USE

16. INFORMATION IN BRAILLE

Tecfidera 120 mg
MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS
HEAT SEALED BLISTER CARD

1. NAME OF THE MEDICINAL PRODUCT

Tecfidera 120 mg Gastro-resistant hard capsules
Dimethyl fumarate

2. NAME OF THE MARKETING AUTHORISATION HOLDER

Biogen Idec

3. EXPIRY DATE

EXP

4. BATCH NUMBER

Lot

5. OTHER

Morning
Evening
Mon.
Tue.
Wed.
Thu.
Fri.
Sat.
Sun.

Sun as a symbol
Moon as a symbol

120 mg capsules
14 gastro-resistant hard capsules

**Oral use**
Each capsule contains 120 mg dimethyl fumarate
Read the package leaflet before use
Keep out of the sight and reach of children

**Do not store above 30ºC**

**Store in original package in order to protect from light**

Medicinal product subject to medical prescription
<table>
<thead>
<tr>
<th>MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>BLISTER FOIL</strong></td>
</tr>
<tr>
<td>1. NAME OF THE MEDICINAL PRODUCT</td>
</tr>
<tr>
<td>Tecfidera 120 mg</td>
</tr>
<tr>
<td>dimethyl fumarate</td>
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<tr>
<td>2. NAME OF THE MARKETING AUTHORISATION HOLDER</td>
</tr>
<tr>
<td>3. EXPIRY DATE</td>
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<tr>
<td>EXP</td>
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<td>4. BATCH NUMBER</td>
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<tr>
<td>Lot</td>
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<tr>
<td>5. OTHER</td>
</tr>
</tbody>
</table>
**PARTICULARS TO APPEAR ON THE OUTER PACKAGING**

**OUTER CARTON**

<table>
<thead>
<tr>
<th>1. NAME OF THE MEDICINAL PRODUCT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tecfidera 240 mg Gastro-resistant hard capsules</td>
</tr>
<tr>
<td>Dimethyl fumarate</td>
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<thead>
<tr>
<th>2. STATEMENT OF ACTIVE SUBSTANCE(S)</th>
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<tbody>
<tr>
<td>Each capsule contains 240 mg dimethyl fumarate</td>
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</table>

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<tr>
<th>3. LIST OF EXCIPIENTS</th>
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</table>

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<tr>
<th>4. PHARMACEUTICAL FORM AND CONTENTS</th>
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</thead>
<tbody>
<tr>
<td>56 gastro-resistant hard capsules</td>
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<tr>
<td>168 gastro-resistant hard capsules</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>5. METHOD AND ROUTE(S) OF ADMINISTRATION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Read the package leaflet before use</td>
</tr>
<tr>
<td>Oral use</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN</th>
</tr>
</thead>
<tbody>
<tr>
<td>Keep out of the sight and reach of children</td>
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<tr>
<th>7. OTHER SPECIAL WARNING(S), IF NECESSARY</th>
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<tr>
<th>8. EXPIRY DATE</th>
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<td>EXP</td>
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<tr>
<th>9. SPECIAL STORAGE CONDITIONS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Do not store above 30ºC</td>
</tr>
<tr>
<td>Store in original package in order to protect from light</td>
</tr>
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<tr>
<th>10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE</th>
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**11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER**

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United Kingdom

**12. MARKETING AUTHORISATION NUMBER(S)**

EU/1/13/837/002
EU/1/13/837/003

**13. BATCH NUMBER**

Lot

**14. GENERAL CLASSIFICATION FOR SUPPLY**

Medicinal product subject to medical prescription

**15. INSTRUCTIONS ON USE**

**16. INFORMATION IN BRAILLE**

Tecfidera 240 mg
### MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS

**HEAT SEALED BLISTER CARD**

<table>
<thead>
<tr>
<th>1. NAME OF THE MEDICINAL PRODUCT</th>
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</thead>
<tbody>
<tr>
<td>Tecfidera 240 mg Gastro-resistant hard capsules</td>
</tr>
<tr>
<td>Dimethyl fumarate</td>
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<tr>
<th>2. NAME OF THE MARKETING AUTHORISATION HOLDER</th>
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<td>Biogen Idec</td>
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<tr>
<th>3. EXPIRY DATE</th>
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<td>EXP</td>
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<table>
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<tr>
<th>4. BATCH NUMBER</th>
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<tr>
<th>5. OTHER</th>
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*Moon as a symbol*
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</tr>
</thead>
<tbody>
<tr>
<td>BLISTER FOIL</td>
</tr>
</tbody>
</table>

1. **NAME OF THE MEDICINAL PRODUCT**

   Tecfidera 240 mg  
   dimethyl fumarate

2. **NAME OF THE MARKETING AUTHORISATION HOLDER**

3. **EXPIRY DATE**

   EXP

4. **BATCH NUMBER**

   Lot

5. **OTHER**
B. PACKAGE LEAFLET
Package leaflet: Information for the patient

Tecfidera 120 mg gastro-resistant hard capsules
Tecfidera 240 mg gastro-resistant hard capsules
dimethyl fumarate

Read all of this leaflet carefully before you start taking this medicine because it contains important information for you.
- Keep this leaflet. You may need to read it again.
- If you have any further questions, ask your doctor or pharmacist.
- This medicine has been prescribed for you only. Do not pass it on to others. It may harm them, even if their signs of illness are the same as yours.
- If you get any side effects, talk to your doctor or pharmacist. This includes any possible side effects not listed in this leaflet. See section 4.

What is in this leaflet
1. What Tecfidera is and what it is used for
2. What you need to know before you take Tecfidera
3. How to take Tecfidera
4. Possible side effects
5. How to store Tecfidera
6. Contents of the pack and other information

1. What Tecfidera is and what it is used for

What Tecfidera is

Tecfidera is a medicine that contains the active substance dimethyl fumarate.

What Tecfidera is used for

Tecfidera is used to treat relapsing-remitting multiple sclerosis (MS).

MS is a long-term condition that affects the central nervous system (CNS), including the brain and the spinal cord. Relapsing-remitting MS is characterised by repeated attacks (relapses) of nervous system symptoms. Symptoms vary from patient to patient but typically include walking difficulties, feeling off balance and visual difficulties. These symptoms may disappear completely when the relapse is over, but some problems may remain.

How Tecfidera works

Tecfidera seems to work by stopping the body’s defence system from damaging your brain and spinal cord. This may also help to delay future worsening of your MS.

2. What you need to know before you take Tecfidera

Do not take Tecfidera:

- if you are allergic to dimethyl fumarate or any of the other ingredients of this medicine (listed in section 6).
Warnings and precautions

Tecfidera may affect your **white blood cell counts**, your **kidneys** and **liver**. Before you start Tecfidera, your doctor will do a blood test to count the number of your white blood cells and will check that your kidneys and liver are working properly. Your doctor will test these periodically during treatment.

**Talk to your doctor** before taking Tecfidera if you have:
- severe **kidney** disease
- severe **liver** disease
- a disease of the **stomach** or **bowel**
- a serious **infection** (such as pneumonia)

Children and adolescents

Tecfidera should **not be used** in children and adolescents below 18 years old. The safety and effectiveness of Tecfidera in this age group are not known.

Other medicines and Tecfidera

**Tell your doctor or pharmacist** if you are taking, have recently taken or might take any medicines, in particular:
- Medicines that contain **fumaric acid esters** (fumarates) used to treat psoriasis.
- Medicines that affect the body’s immune system including **other medicines used to treat** MS, such as fingolimod, natalizumab or mitoxantrone or some commonly used **cancer treatments**.
- Medicines that affect the kidneys including **some antibiotics** (used to treat infections), **“water tablets”** (**diuretics**), **certain types of painkillers** (such as ibuprofen and other similar anti-inflammatories and medicines purchased without a doctor’s prescription) and medicines that contain **lithium**.
- **Oral contraceptives** (also called “hormonal contraceptives” or “**the pill**”). Tecfidera may make oral contraceptives less effective. Use an extra form of contraception (such as a condom) while you’re taking Tecfidera.
- **Vaccinations** given while taking Tecfidera may be less effective than normal. Taking Tecfidera with certain types of vaccine (**live vaccines**) may cause you to get an infection and should therefore be avoided.

Tecfidera with food and alcohol

Consumption of more than a small quantity (more than 50 ml) of strong alcoholic drinks (more than 30% alcohol by volume, e.g. spirits) should be avoided within an hour of taking Tecfidera, as alcohol can interact with this medicine. This could cause inflammation of the stomach (**gastritis**), especially in people already prone to gastritis.

Pregnancy and breast-feeding

If you are pregnant or breast-feeding, think you may be pregnant or are planning to have a baby, ask your doctor or pharmacist for advice before taking this medicine.

**Pregnancy**

Do not use Tecfidera if you are pregnant unless you have discussed this with your doctor.

**Breast-feeding**

It is not known whether the ingredients in Tecfidera pass into breast milk. Tecfidera is not to be used during breast-feeding. Your doctor will help you decide whether you should stop breast-feeding, or
stop using Tecfidera. This involves balancing the benefit of breast-feeding for your child, and the benefit of therapy for you.

**Driving and using machines**

The effect of Tecfidera on the ability to drive or use machines is not known. Your doctor will tell you whether your illness allows you to drive vehicles and use machines safely.

3. **How to take Tecfidera**

Always take this medicine exactly as your doctor has told you. Check with your doctor if you are not sure.

**Starting dose**

120 mg twice a day.  
Take this starting dose for the first 7 days, then take the regular dose.

**Regular dose**

240 mg twice a day.

Swallow each capsule whole, with some water. Do not divide, crush, dissolve, suck or chew the capsule as this may increase some side effects.

**Take Tecfidera with food** – it may help to reduce some of the very common side effects (listed in Section 4)

**If you take more Tecfidera than you should**

If you have taken too many capsules, *talk to your doctor straight away*.

**If you forget to take Tecfidera**

If you forget or miss a dose, *do not take a double dose*.  
You may take the missed dose if you leave at least 4 hours between the doses. Otherwise wait until your next planned dose.

If you have any further questions on the use of this medicine, ask your doctor or pharmacist.

4. **Possible side effects**

Like all medicines, this medicine can cause side effects, although not everybody gets them.

**Serious effects**

Allergic reactions - these are uncommon and may affect *up to 1 in 100 people*

Reddening of the face or body (*flushing*) is a very common (*may affect more than 1 in 10 people*) side effect. However, if you become flushed and get any of these signs:

- swelling of the face, lips, mouth or tongue
- wheezing, difficulty breathing or shortness of breath
Stop taking Tecfidera and call a doctor straight away.

**Very common side effects**

These may affect *more than 1 in 10 people*:
- reddening of the face or body feeling warm, hot, burning or itchy (*flushing*)
- loose stools (*diarrhoea*)
- feeling sick (*nausea*)
- stomach pain or stomach cramps

**taking your medicine with food** can help to reduce the side effects above.

Substances called ketones, which are naturally produced in the body, very commonly show up in urine tests while taking Tecfidera.

**Talk to your doctor** about how to manage these side effects. Your doctor may reduce your dose. Do not reduce your dose unless your doctor tells you to.

**Common side effects**

These may affect *up to 1 in 10 people*:
- inflammation of the lining of the intestines (*gastroenteritis*)
- being sick (*vomiting*)
- indigestion (*dyspepsia*)
- inflammation of the lining of the stomach (*gastritis*)
- gastrointestinal disorder
- burning sensation
- hot flush, feeling hot
- itchy skin (*pruritus*)
- rash
- pink or red blotches on the skin (*erythema*)

**Common side effects, which may show up in your blood or urine tests**

- low levels of white blood cells (*lymphopenia, leucopenia*) in the blood. Reduced white blood cells could mean your body is less able to fight an infection. If you have a serious infection (such as pneumonia), talk to your doctor immediately.
- proteins (*albumin*) in urine
- increase in levels of liver enzymes (*ALT, AST*) in the blood

**Reporting of side effects**

If you get any side effects, talk to your doctor or, pharmacist. This includes any possible side effects not listed in this leaflet. You can also report side effects directly via the [national reporting system](#) listed in Appendix V. By reporting side effects you can help provide more information on the safety of this medicine.

5. **How to store Tecfidera**

Keep this medicine out of the sight and reach of children.

Do not use this medicine after the expiry date which is stated on the carton after “EXP”.

The expiry date refers to the last day of that month.

Do not store above 30°C.

Store in the original package in order to protect from light.
Do not throw away any medicines via wastewater or household waste. Ask your pharmacist how to dispose of medicines you no longer use. These measures will help protect the environment.

6. Contents of the pack and other information

What Tecfidera contains

The active substance is dimethyl fumarate.
Tecfidera 120 mg: Each capsule contains 120 mg of dimethyl fumarate.
Tecfidera 240 mg: Each capsule contains 240 mg of dimethyl fumarate.

The other ingredients are microcrystalline cellulose, croscarmellose sodium, tale, silica, hydrophobic colloidal, magnesium stearate, triethyl citrate, methacrylic acid – methyl methacrylate copolymer (1:1), methacrylic acid – ethyl acrylate copolymer (1:1) dispersion 30%, simeticone, sodium laurilsulfate, polysorbate 80, gelatin, titanium dioxide (E171), brilliant blue FCF (E133), yellow iron oxide (E172), shellac, potassium hydroxide and black iron oxide (E172).

What Tecfidera looks like and contents of the pack

Tecfidera 120 mg gastro-resistant hard capsules are green and white and printed with ‘BG-12 120 mg’ and are available in packs containing 14 capsules.

Tecfidera 240 mg gastro-resistant hard capsules are green and printed with ‘BG-12 240 mg’ and are available in packs containing 56 or 168 capsules.

Not all pack sizes may be marketed.

Marketing Authorisation Holder

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Manufacturer

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This leaflet was last revised in {MM YYY}.

Detailed information on this medicine is available on the European Medicines Agency web site: http://www.ema.europa.eu.