ANNEX I SUMMARY OF PRODUCT CHARACTERISTICS

This medicinal product is subject to additional monitoring. This will allow quick identification of new safety information. Healthcare professionals are asked to report any suspected adverse reactions. See section 4.8 for how to report adverse reactions.

1. NAME OF THE MEDICINAL PRODUCT

Litfulo 50 mg hard capsules

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each hard capsule contains ritlecitinib tosylate equivalent to 50 mg ritlecitinib.

Excipient(s) with known effect

Each hard capsule contains 21.27 mg of lactose monohydrate.

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Hard capsule (capsule)

Opaque hard capsules, yellow body and blue cap approximately 16 mm long and 6 mm wide, of which the body is printed with "RCB 50" and the cap is printed with "Pfizer" in black.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Litfulo is indicated for the treatment of severe alopecia areata in adults and adolescents 12 years of age and older (see section 5.1).

4.2 Posology and method of administration

Treatment should be initiated and supervised by a healthcare professional experienced in the diagnosis and treatment of alopecia areata.

Posology

The recommended dose is 50 mg once daily.

The benefit-risk of treatment should be re-assessed at regular intervals on an individual basis.

Consideration should be given to discontinuing patients who show no evidence of therapeutic benefit after 36 weeks.

Table 1. Laboratory measures and monitoring guidance

Laboratory	Monitoring guidance	Action
measures		
Platelet count		Treatment should be
	Before treatment initiation, 4 weeks after	discontinued if platelet count is
	initiation, and thereafter according to	$< 50 \times 10^{3} / \text{mm}^{3}$.
Lymphocytes	routine patient management.	Treatment should be interrupted
		if ALC is $< 0.5 \times 10^3 / \text{mm}^3$ and
		may be restarted once ALC
		return above this value.

Abbreviation: ALC = absolute lymphocyte count

Treatment initiation

Treatment with ritlecitinib should not be initiated in patients with an absolute lymphocyte count (ALC) $< 0.5 \times 10^3 / \text{mm}^3$ or a platelet count $< 100 \times 10^3 / \text{mm}^3$ (see section 4.4).

Treatment interruption or discontinuation

If a patient develops a serious infection or opportunistic infection, ritlecitinib should be interrupted until the infection is controlled (see section 4.4).

Interruption or discontinuation of treatment may be needed for management of haematologic abnormalities as described in Table 1.

If treatment interruption is needed, the risk of significant loss of regrown scalp hair after a temporary treatment interruption for less than 6 weeks is low.

Missed doses

If a dose is missed, patients should be advised to take the dose as soon as possible unless it is less than 8 hours before the next dose, in which case the patient should not take the missed dose. Thereafter, dosing should be resumed at the regular scheduled time.

Special populations

Renal impairment

No dose adjustment is required in patients with mild, moderate, or severe renal impairment (see section 5.2).

Ritlecitinib has not been studied in patients with end-stage renal disease (ESRD) or in patients with renal transplants and is therefore not recommended for use in these patients.

Hepatic impairment

No dose adjustment is required in patients with mild (Child Pugh A) or moderate (Child Pugh B) hepatic impairment (see section 5.2). Ritlecitinib is contraindicated in patients with severe (Child Pugh C) hepatic impairment (see section 4.3).

Elderly

No dose adjustment is required for patients \geq 65 years of age. There are limited data in patients \geq 65 years of age.

Paediatric population

No dose adjustment is required for adolescents 12 to < 18 years of age.

The safety and efficacy of Litfulo in children under 12 years of age have not yet been established. No data are available.

Method of administration

Oral use.

Litfulo is to be taken once daily with or without food.

Capsules should be swallowed whole and should not be crushed, split or chewed, because these methods of administration have not been studied in clinical trials.

4.3 Contraindications

- Hypersensitivity to the active substance(s) or to any of the excipients listed in section 6.1.
- Active serious infections, including tuberculosis (TB) (see section 4.4).
- Severe hepatic impairment (see section 4.2).
- Pregnancy and breast-feeding (see section 4.6).

4.4 Special warnings and precautions for use

Serious infections

Serious infections have been reported in patients receiving ritlecitinib. The most frequent serious infections have been appendicitis, COVID-19 infection (including pneumonia), and sepsis. Treatment with ritlecitinib must not be initiated in patients with an active, serious infection (see section 4.3).

The risks and benefits of treatment should be considered in patients:

- with chronic or recurrent infection
- who have been exposed to tuberculosis (TB)
- with a history of serious or an opportunistic infection
- who have resided or traveled in areas of endemic TB or mycoses, or
- with underlying conditions that may predispose them to infection

Patients should be closely monitored for the development of signs and symptoms of infection during and after treatment with ritlecitinib. Treatment should be interrupted if a patient develops a serious or opportunistic infection. A patient who develops a new infection during treatment with ritlecitinib should undergo prompt and complete diagnostic testing appropriate for an immunocompromised patient, appropriate antimicrobial therapy should be initiated, and the patient should be closely monitored. If interrupted, ritlecitinib may be resumed once the infection is controlled.

As there is a higher incidence of infections in elderly and in the diabetic population in general, caution should be exercised when treating the elderly and patients with diabetes, and particular attention paid with respect to occurrence of infections.

Tuberculosis

Patients should be screened for TB before starting therapy with ritlecitinib. Ritlecitinib must not be given to patients with active TB (see section 4.3). Anti-TB therapy should be started prior to initiating therapy with ritlecitinib in patients with a new diagnosis of latent TB or previously untreated latent TB. In patients with a negative latent TB test, anti-TB therapy should still be considered before initiating treatment with ritlecitinib in those at high risk and screening for patients at high risk for TB during treatment with ritlecitinib should be considered.

Viral reactivation

Viral reactivations, including cases of herpes virus reactivation (e.g., herpes zoster), have been reported (see section 4.8). If a patient develops herpes zoster, temporary interruption of treatment may be considered until the episode resolves.

Screening for viral hepatitis should be performed in accordance with clinical guidelines before starting therapy with ritlecitinib. Patients with evidence of hepatitis B or C infection were excluded from studies with ritlecitinib. Monitoring for reactivation of viral hepatitis according to clinical guidelines is recommended during ritlecitinib treatment. If there is evidence of reactivation, a liver specialist should be consulted.

Malignancy (including non-melanoma skin cancer)

Malignancies, including non-melanoma skin cancer (NMSC) have been reported in patients receiving ritlecitinib.

It is not known whether selective JAK3 inhibition may be associated with adverse reactions of Janus Kinase (JAK) inhibition predominantly involving JAK1 and JAK2. In a large randomised active-controlled study of tofacitinib (another JAK inhibitor) in rheumatoid arthritis (RA) patients 50 years and older with at least one additional cardiovascular risk factor, a higher rate of malignancies, particularly lung cancer, lymphoma and NMSC, was observed with tofacitinib compared to tumour necrosis factor (TNF) inhibitors.

Limited clinical data are available to assess the potential relationship of exposure to ritlecitinib and the development of malignancies. Long-term safety evaluations are ongoing. The risks and benefits of ritlecitinib treatment should be considered prior to initiating or continuing therapy in patients with a known malignancy other than a successfully treated NMSC or cervical cancer.

Periodic skin examination is recommended for patients who are at increased risk of skin cancer.

<u>Major adverse cardiovascular events (MACE), deep venous thrombosis (DVT) and pulmonary</u> embolism (PE)

Events of venous and arterial thromboembolism, including MACE, have been reported in patients receiving ritlecitinib.

It is not known whether selective JAK3 inhibition may be associated with adverse reactions of JAK inhibition predominantly involving JAK1 and JAK2. In a large randomised active-controlled study of tofacitinib (another JAK inhibitor) in RA patients 50 years and older with at least one additional cardiovascular risk factor, a higher rate of MACE, defined as cardiovascular death, non-fatal myocardial infarction and non-fatal stroke, and a dose-dependent higher rate of venous thromboembolism including DVT and PE were observed with tofacitinib compared to TNF inhibitors.

Long-term safety evaluations for ritlecitinib are ongoing. Ritlecitinib should be used with caution in patients with known risk factors for thromboembolism. In patients with a suspected thromboembolic event, discontinuation of ritlecitinib and prompt re-evaluation is recommended. The risks and benefits of ritlecitinib treatment should be considered prior to initiating therapy in patients.

Neurological events

Ritlecitinib-related axonal dystrophy has been observed in chronic Beagle dog toxicity studies (see section 5.3). Treatment with ritlecitinib should be discontinued in case unexplained neurological symptoms occur.

Haematologic abnormalities

Treatment with ritlecitinib was associated with decreases in lymphocytes and platelets (see section 4.8). Prior to initiating treatment with ritlecitinib, ALC and platelet counts should be performed. Treatment with ritlecitinib should not be initiated in patients with an ALC $< 0.5 \times 10^3 / \text{mm}^3$ or a platelet count $< 100 \times 10^3 / \text{mm}^3$. After initiating treatment with ritlecitinib, treatment interruption or discontinuation are recommended based on ALC and platelet count abnormalities (see section 4.2).

ALC and platelet counts are recommended at 4 weeks after initiation of therapy with ritlecitinib, and thereafter according to routine patient management.

Vaccinations

No data are available on the response to vaccination in patients receiving ritlecitinib. Use of live attenuated vaccines should be avoided during or immediately prior to ritlecitinib treatment. Prior to initiating ritlecitinib, it is recommended that patients are brought up to date with all immunisations, including prophylactic herpes zoster vaccinations, in agreement with current immunisation guidelines.

Elderly

There are limited data in patients \geq 65 years of age. Age appeared to be a risk factor for lower ALC in patients \geq 65 years of age.

Excipients with known effect

Lactose

Patients with rare hereditary problems of galactose intolerance, total lactase deficiency or glucose-galactose malabsorption should not take this medicinal product.

4.5 Interaction with other medicinal products and other forms of interaction

Potential for other medicinal products to affect the pharmacokinetics of ritlecitinib

The coadministration of multiple 200 mg doses of itraconazole, a strong CYP3A inhibitor, increased the area under curve (AUC)_{inf} of ritlecitinib by approximately 15%. This is not considered clinically significant and, therefore dose adjustment is not required when ritlecitinib is coadministered with CYP3A inhibitors.

The coadministration of multiple 600 mg doses of rifampicin, a strong inducer of CYP enzymes, decreased the AUC_{inf} of ritlecitinib by approximately 44%. This is not considered clinically significant and, therefore dose adjustment is not required when ritlecitinib is coadministered with inducers of CYP enzymes.

Potential for ritlecitinib to affect the pharmacokinetics of other medicinal products

Multiple doses of 200 mg once daily ritlecitinib increased the AUC_{inf} and C_{max} of midazolam a CYP3A4 substrate, by approximately 2.7-fold and 1.8-fold, respectively. Ritlecitinib is a moderate inhibitor of CYP3A; caution should be exercised with concomitant use of ritlecitinib with CYP3A substrates (e.g., quinidine, cyclosporine, dihydroergotamine, ergotamine, pimozide) where moderate concentration changes may lead to serious adverse reactions. Dose adjustment recommendations for the CYP3A substrate (e.g., colchicine, everolimus, tacrolimus, sirolimus) should be considered.

Multiple doses of 200 mg once daily ritlecitinib increased the AUC_{inf} and C_{max} of caffeine, a CYP1A2 substrate, by approximately 2.7-fold and 1.1-fold, respectively. Ritlecitinib is a moderate inhibitor of CYP1A2; caution should be exercised with concomitant use of ritlecitinib with other CYP1A2 substrates (e.g., tizanidine) where moderate concentration changes may lead to serious adverse reactions. Dose adjustment recommendations for the CYP1A2 substrate (e.g., theophylline, pirfenidone) should be considered.

The coadministration of a single 400 mg dose of ritlecitinib increased the AUC_{inf} of sumatriptan (an organic cation transporter [OCT]1 substrate) by approximately 1.3 to 1.5-fold relative to sumatriptan dose given alone. The increase in sumatriptan exposure is not considered clinically relevant. Caution should be exercised with concomitant use of ritlecitinib with OCT1 substrates where small concentration changes may lead to serious adverse reactions.

Ritlecitinib did not produce clinically significant changes in the exposures of oral contraceptives (e.g., ethinyl oestradiol or levonorgestrel), CYP2B6 substrates (e.g., efavirenz), CYP2C substrates (e.g., tolbutamide), or substrates of organic anion transporter (OAT)P1B1, breast cancer resistant protein (BCRP), and OAT3 (e.g., rosuvastatin).

Paediatric population

Interaction studies have only been performed in adults.

4.6 Fertility, pregnancy and lactation

Women of childbearing potential

Ritlecitinib is not recommended in women of childbearing potential not using contraception. Women of childbearing potential have to use effective contraception during treatment and for 1 month following the final dose of Litfulo.

Pregnancy

There are no or limited data from the use of ritlecitinib in pregnant women. Studies in animals have shown reproductive toxicity (see section 5.3). Ritlecitinib was teratogenic in rats and rabbits at high doses (see section 5.3). Litfulo is contraindicated during pregnancy (see section 4.3).

Breast-feeding

Available pharmacodynamic/toxicological data in animals have shown excretion of ritlecitinib in milk (see section 5.3). A risk to newborns/infants cannot be excluded. Litfulo is contraindicated during breast-feeding (see section 4.3).

Fertility

The effect of ritlecitinib on human fertility has not been evaluated. There were no effects on fertility in rats at clinically relevant exposures (see section 5.3).

4.7 Effects on ability to drive and use machines

Litfulo has no or negligible influence on the ability to drive and use machines.

4.8 Undesirable effects

Summary of the safety profile

The most frequently reported adverse reactions are diarrhoea (9.2%), acne (6.2%), upper respiratory tract infections (6.2%), urticaria (4.6%), rash (3.8%), folliculitis (3.1%), and dizziness (2.3%).

Tabulated list of adverse reactions

A total of 1630 patients were treated with ritlecitinib in placebo-controlled studies of alopecia areata representing 2303 patient-years of exposure. Three placebo-controlled studies were integrated (130 participants on 50 mg daily and 213 participants on placebo) to evaluate the safety of ritlecitinib in comparison to placebo for up to 24 weeks after treatment initiation.

Table 2 lists all adverse reactions observed in alopecia areata placebo-controlled studies presented by system organ class and frequency, using the following categories: very common ($\geq 1/10$); common ($\geq 1/100$ to < 1/10); uncommon ($\geq 1/1000$); rare ($\geq 1/10000$) to < 1/1000); very rare (< 1/10000). Within each frequency grouping, adverse reactions are presented in order of decreasing seriousness.

Table 2. Adverse reactions

System organ class	Common	Uncommon
Infections and infestations	Herpes zoster	
	Folliculitis	
	Upper respiratory tract	
	infections	
Nervous system disorders	Dizziness	
Gastrointestinal disorders	Diarrhoea	
Skin and subcutaneous tissue	Acne	
disorders	Urticaria	
	Rash	
Investigations	Blood creatine phosphokinase	Platelet count decreased
	increased	Lymphocyte count decreased
		Alanine aminotransferase
		increased $> 3 \times ULN^a$
		Aspartate aminotransferase
		increased $> 3 \times ULN^a$

a. Includes changes detected during laboratory monitoring

Description of selected adverse reactions

Infections

In the placebo-controlled studies, for up to 24 weeks, overall infections have been reported in 31% of patients (80.35 per 100 patient-years) treated with placebo and 33% of patients (74.53 per 100 patient-years) treated with ritlecitinib 50 mg. In study AA-I, for up to 48 weeks, overall infections were reported in 51% of patients (89.32 per 100 patient-years) treated with ritlecitinib 50 mg or higher.

Among all patients treated with ritlecitinib in the integrated safety analysis, including the long-term study and a study in vitiligo, overall infections were reported in 45.4% of patients (50.02 per 100 patient-years) treated with ritlecitinib 50 mg or higher. Most infections were mild or moderate in severity.

In the placebo-controlled studies the percentage of patients reporting infection-related adverse reaction of herpes zoster were 1.5% in the ritlecitinib 50 mg group compared to 0 in placebo. All herpes zoster events were non-serious; 1 patient receiving ritlecitinib 200/50 mg (200 mg once daily for 4 weeks followed by 50 mg once daily) experienced an event of varicella zoster virus infection that met criteria as an opportunistic infection (multi-dermatomal herpes zoster). In study AA-I, for up to 48 weeks, 2.3% of patients (2.61 per 100 patient-years) treated with ritlecitinib 50 mg or higher reported herpes zoster Among all patients treated with ritlecitinib in the integrated safety analysis, including the long-term study and a study in vitiligo, the rate of herpes zoster was 1.10 per 100 patient-years in patients treated with ritlecitinib 50 mg or higher.

In the placebo-controlled studies, for up to 24 weeks, no serious infections were reported in patients treated with placebo or ritlecitinib 50 mg. The proportion and rate of serious infections in patients treated with ritlecitinib 200/50 mg was 0.9% (2.66 per 100 patient-years). In study AA-I, for up to 48 weeks, serious infections were reported in 0.8% of patients (0.86 per 100 patient-years) treated with ritlecitinib 50 mg or higher. Among all patients treated with ritlecitinib in the integrated safety analysis, including the long-term study and a study in vitiligo, the proportion and rate of serious infection in ritlecitinib 50 mg or higher was 0.8% (0.59 per 100 patient-years).

Opportunistic infections

Opportunistic infections of multi-dermatomal herpes zoster were reported in 1 patient (0.50 per 100 patient-years) treated with ritlecitinib 200/50 mg in the placebo-controlled studies, no patients in study AA-I, for up to 48 weeks, and 2 patients (0.09 per 100 patient-years) treated with ritlecitinib 50 mg

or higher in the integrated safety analysis, including the long-term study and a study in vitiligo. Cases of opportunistic herpes zoster were mild or moderate in severity.

Decreased lymphocyte count

In the placebo-controlled studies, for up to 24 weeks, and study AA-I, for up to 48 weeks, treatment with ritlecitinib was associated with a decrease in lymphocyte count. Maximum effects on lymphocytes were observed within 4 weeks, after which lymphocyte count remained stable at a lower level with continued therapy. Among all patients treated with ritlecitinib in the integrated safety analysis, including the long-term study and a study in vitiligo, confirmed ALC $< 0.5 \times 10^3 / \text{mm}^3$ occurred in 2 participants (< 0.1%) treated with ritlecitinib 50 mg.

Decreased platelet count

In the placebo-controlled studies, for up to 24 weeks, and study AA-I, for up to 48 weeks, treatment with ritlecitinib was associated with a decrease in platelet count. Maximum effects on platelets were observed within 4 weeks, after which platelet count remained stable at a lower level with continued therapy. Among all patients treated with ritlecitinib in the integrated safety analysis, including the long-term study and a study in vitiligo, 1 patient (< 0.1%) treated with ritlecitinib 50 mg or higher had a confirmed platelet count $< 100 \times 10^3/\text{mm}^3$.

Creatine phosphokinase (CPK) elevations

In the placebo-controlled studies, for up to 24 weeks, events of blood CPK increased were reported in 2 patients (1.5%) treated with ritlecitinib 50 mg. In study AA-I, for up to 48 weeks, events of blood CPK increased were reported in 3.8% of patients treated with ritlecitinib 50 mg or higher. CPK elevations >5x upper limit of normal (ULN) were reported in 2 (0.9%) of patients treated with placebo and 5 (3.9%) of patients treated with ritlecitinib 50 mg. In study AA-I, for up to 48 weeks, CPK elevations >5x ULN were reported in 6.6% of patients treated with ritlecitinib 50 mg or higher. Most elevations were transient and none led to discontinuation.

Increased transaminases

In the placebo-controlled studies, for up to 24 weeks, events of increases in ALT and AST values $(>3 \times ULN)$ were reported in 3 patients (0.9%) and 2 patients (0.6%) treated with ritlecitinib 50 mg or higher, respectively. Most elevations were transient, and none led to discontinuation.

Paediatric population

A total of 181 adolescents (12 to < 18 years of age) were enrolled in ritlecitinib alopecia areata placebo-controlled studies.

The safety profile observed in adolescents was similar to that of the adult population.

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

Ritlecitinib was administered in placebo-controlled studies up to a single oral dose of 800 mg and multiple oral doses of 400 mg daily for 14 days. No specific toxicities were identified. In case of overdose, it is recommended that the patient be monitored for signs and symptoms of adverse reactions (see section 4.8). There is no specific antidote for overdose with ritlecitinib. Treatment should be symptomatic and supportive.

Pharmacokinetics (PK) data up to and including a single oral dose of 800 mg in healthy adult volunteers indicate that more than 90% of the administered dose is expected to be eliminated within 48 hours.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Immunosuppressants, Janus-associated kinase (JAK) inhibitors, ATC code: L04AF08

Mechanism of action

Ritlecitinib irreversibly and selectively inhibits Janus kinase (JAK) 3 and the tyrosine kinase expressed in hepatocellular carcinoma (TEC) family by blocking the adenosine triphosphate (ATP) binding site. In cellular settings, ritlecitinib specifically inhibits γ -common cytokines (IL-2, IL-4, IL-7, IL-15 and IL-21) signalling through JAK3-dependent common- γ chain receptors. Additionally, ritlecitinib inhibits TEC family of kinases, resulting in reduced cytolytic activity of NK cells and CD8+ T cells.

JAK3 and TEC family mediated signalling pathways are both involved in alopecia areata pathogenesis, although complete pathophysiology is still not understood.

Pharmacodynamic effects

Lymphocyte subsets

In patients with alopecia areata, treatment with ritlecitinib was associated with dose-dependent early decreases in absolute lymphocyte levels, T lymphocytes (CD3) and T lymphocyte subsets (CD4 and CD8). After the initial decrease, the levels partially recovered and remained stable up to 48 weeks. There was no change observed in B lymphocytes (CD19) in any treatment group. There was a dose-dependent early decrease in NK cells (CD16/56) which remained stable at the lower level up to Week 48.

Immunoglobulins

In patients with alopecia areata, treatment with ritlecitinib was not associated with clinically meaningful changes in Immunoglobulin (Ig)G, IgM or IgA up to Week 48, indicating a lack of systemic humoral immunosuppression.

Clinical efficacy and safety

The efficacy and safety of ritlecitinib was evaluated in a pivotal, randomised, double-blind, placebo-controlled study (study AA-I) in alopecia areata patients 12 years of age and older with $\geq 50\%$ scalp hair loss, including alopecia totalis and alopecia universalis. The dose-response of ritlecitinib was also evaluated in this study. The study treatment period consisted of a placebo-controlled 24-week period and a 24-week extension period. Study AA-I evaluated a total of 718 patients who were randomised to one of the following treatment regimens for 48 weeks: 1) 200 mg once daily for 4 weeks followed by 50 mg once daily for 44 weeks; 2) 200 mg once daily for 4 weeks followed by 30 mg once daily for 44 weeks; 3) 50 mg once daily for 48 weeks; 4) 30 mg once daily for 48 weeks; 5) 10 mg once daily for 48 weeks; 6) placebo for 24 weeks followed by 200 mg once daily for 4 weeks and 50 mg once daily for 20 weeks; or 7) placebo for 24 weeks followed by 50 mg for 24 weeks.

This study assessed as primary outcome the proportion of subjects who achieved a SALT (Severity of Alopecia Tool) score of ≤ 10 (90% or more scalp hair coverage) at Week 24. Additionally, this study assessed as key secondary outcome the Patient's Global Impression of Change (PGI-C) response at Week 24 and also assessed as secondary outcomes SALT score of ≤ 20 (80% or more scalp hair coverage) at Week 24 and improvements in regrowth of eyebrows and/or eyelashes at Week 24.

Baseline characteristics

Male or female patients 12 years of age and older, were assessed in study AA-I. All patients had alopecia areata with $\geq 50\%$ scalp hair loss (SALT [Severity of Alopecia Tool] score ≥ 50) without evidence of terminal hair regrowth within the previous 6 months and with the current episode of scalp hair loss ≤ 10 years and no other known cause of hair loss (e.g., androgenetic alopecia).

Across all treatment groups 62.1% were female, 68.0% were White, 25.9% were Asian, and 3.8% were Black or African American. The mean age of patients was 33.7 years and the majority (85.4%) were adults (≥ 18 years of age). A total of 105 (14.6%) patients 12 to < 18 years of age and 20 (2.8%) patients 65 years of age and older were enrolled. The mean (SD) baseline absolute SALT score ranged from 88.3 (16.87) to 93.0 (11.50) across treatment groups; among patients without alopecia totalis/alopecia universalis at baseline, the mean SALT score ranged from 78.3 to 87.0. The majority of patients had abnormal eyebrows (83.0%) and eyelashes (74.7%) at baseline across treatment groups. The median duration since alopecia areata diagnosis was 6.9 years and the median duration of the current alopecia areata episode was 2.5 years. Randomisation was stratified by alopecia totalis/alopecia universalis status with 46% of patients classified as alopecia totalis/alopecia universalis based upon a baseline SALT score of 100.

Clinical response

A significantly greater proportion of patients achieved SALT \leq 10 response with ritlecitinib 50 mg compared to placebo at Week 24 (Table 3). The SALT \leq 10 response rate for ritlecitinib 50 mg increased further at Week 48 (Figure 1).

A significantly greater proportion of patients achieved Patient's Global Impression of Change (PGI-C) response with ritlecitinib 50 mg compared to placebo at Week 24 (Table 3) with response rates continuing to increase through Week 48 (Figure 1).

A significantly greater proportion of patients achieved a SALT \leq 20 response with ritlecitinib 50 mg compared to placebo at Week 24 (Table 3). The SALT \leq 20 response rate increased further at Week 48.

Improvements in regrowth of eyebrows and/or eyelashes were seen at Week 24 (Table 3) with ritlecitinib 50 mg among patients with abnormal eyebrows and/or eyelashes at baseline with further increases seen at Week 48.

Treatment effects at Week 24 in subgroups (age, gender, race, region, weight, duration of disease since diagnosis, duration of current episode, prior pharmacologic treatment) were consistent with the results in the overall study population. Treatment effects at Week 24 in the alopecia totalis/alopecia universalis subgroup were lower compared to the non-alopecia totalis/non-alopecia universalis subgroup. Treatment effects at Week 24 in adolescents 12 to less than 18 years of age were consistent with the results in the overall study population.

Table 3. Efficacy results of ritlecitinib at week 24

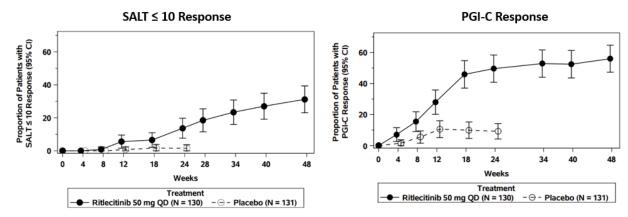
Endpoint	Ritlecitinib 50 mg once daily (N = 130) % Responders	Placebo (N = 131) % Responders	Difference from placebo (95% CI)
$SALT \le 10 \text{ response}^{a,b}$	13.4	1.5	11.9 (5.4, 18.3)
PGI-C response ^{b,c}	49.2	9.2	40.0 (28.9, 51.1)
$SALT \le 20 \text{ response}^{d,e}$	23.0	1.6	21.4 (13.4, 29.5)
EBA response ^f	29.0	4.7	24.3 (14.8, 34.5)
ELA response ^g	28.9	5.2	23.7

		(13.6.34.5)
		(13.0, 34.3)

Abbreviations: EBA = eyebrow assessment; ELA = eyelash assessment; CI = confidence interval; N = total number of patients; PGI-C = Patient's Global Impression of Change; SALT = Severity of Alopecia Tool

- a. SALT \leq 10 responders were patients with scalp hair loss of \leq 10%. SALT scores range from 0 to 100 with 0 = no scalp hair loss and 100 = total scalp hair loss.
- b. Statistically significant with adjustment for multiplicity.
- c. PGI-C responders were patients with a score of "moderately improved" or "greatly improved" based upon a 7-point scale from "greatly improved" to "greatly worsened".
- d. SALT \leq 20 responders were patients with scalp hair loss of \leq 20%. SALT scores range from 0 to 100 with 0 = no scalp hair loss and 100 = total scalp hair loss.
- e. Statistically significant.
- f. EBA response is defined as at least a 2-grade improvement from baseline or normal EBA score on a 4-point scale in patients with abnormal eyebrows at baseline.
- g. ELA response is defined as at least a 2-grade improvement from baseline or normal ELA score on a 4-point scale in patients with abnormal eyelashes at baseline.

Figure 1. SALT \leq 10 and PGI-C response through Week 48



Abbreviations: CI = confidence interval; N = total number of patients; PGI-C = Patient Global Impression of Change; QD = once daily; SALT = Severity of Alopecia Tool

Paediatric population

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

5.2 Pharmacokinetic properties

Absorption

The absolute oral bioavailability of ritlecitinib is about 64%. Based on oral and intravenous administration of the labelled active substance, the relative urinary recovery (oral/intravenous) of labelled compounds was about 89%, indicating a high fraction absorbed (f_a). Peak plasma concentrations are reached within 1 hour following multiple oral doses. Food does not have a clinically significant impact on the extent of ritlecitinib absorption, as a high-fat meal decreased the ritlecitinib C_{max} by ~ 32% and increased AUC_{inf} by ~11%. In placebo-controlled studies, ritlecitinib was administered without regard to meals (see section 4.2).

In vitro, ritlecitinib is a substrate of P-glycoprotein (P-gp) and BCRP. However, as ritlecitinib has a high fraction absorbed (f_a) with both C_{max} and AUC increases in a dose proportional manner (20-200 mg single dose range), P-gp and BCRP are not expected to have a meaningful impact on the absorption of ritlecitinib.

Distribution

After intravenous administration, the volume of distribution of ritlecitinib is about 74 L. Approximately 14% of circulating ritlecitinib is bound to plasma proteins, primarily albumin. The blood/plasma distribution ratio of ritlecitinib is 1.62. Ritlecitinib is a covalent inhibitor that has been shown to bind to off-target proteins such as MAP2K7, DOCK10, albumin, CYP1A2, CYP3A, UGT1A1, and UGT1A4, some of which may have clinical relevance in drug interactions (see section 4.5).

Biotransformation

The metabolism of ritlecitinib is mediated by multiple isoforms of Glutathione S-transferase (GST: cytosolic GST A1/3, M1/3/5, P1, S1, T2, Z1, and microsomal Membrane Associated Proteins involved in Eicosanoid and Glutathione metabolism [MAPEG]1/2/3) and CYP enzymes (CYP3A, CYP2C8, CYP1A2, and CYP2C9), with no single clearance route contributing more than 25%. Hence, medicinal products inhibiting a selective metabolic pathway are unlikely to impact the systemic exposures of ritlecitinib. Specific inhibitors of transporters are unlikely to result in clinically relevant changes in the bioavailability of ritlecitinib.

In a human radiolabeled study, ritlecitinib was the most prevalent circulating species (30.4% of circulating radioactivity) after oral administration, with a major cysteine conjugate metabolite M2 (16.5%), which is pharmacologically inactive.

Elimination

Ritlecitinib is eliminated primarily by metabolic clearance mechanisms, with approximately 4% of the dose excreted as unchanged active substance in urine. Approximately 66% of radiolabeled ritlecitinib dose is excreted in the urine and 20% in the faeces. Following multiple oral doses, steady state was reached approximately by Day 4 due to non-stationary PK. The steady state PK parameters of AUC $_{tau}$ and C_{max} appeared to increase in an approximately dose-proportional manner up to 200 mg with the mean terminal half-life ranging from 1.3 to 2.3 hours.

Special populations

Body weight, gender, genotype, race and age Body weight, gender, GST P1, M1, and T1 genotype, race and age did not have a clinically meaningful effect on ritlecitinib exposure. Adolescents ($\geq 12 \text{ to} < 18 \text{ years}$)

Based on population PK analysis, there was no clinically relevant difference in ritlecitinib exposures in adolescent patients compared to adults.

Paediatric (< 12 years)

The PK of ritlecitinib in children under 12 years of age have not yet been established.

Renal impairment

The AUC $_{24}$ and C $_{max}$ in patients with severe renal impairment (estimated glomerular filtration rate [eGFR] < 30 mL/min) was about 55% and 44% higher, respectively, compared with matched participants with normal renal functions. This was confirmed by popPK analysis. These differences are not considered clinically significant. Ritlecitinib was not studied in patients with mild (eGFR 60 to < 90 mL/min) or moderate (eGFR 30 to < 60 mL/min) renal impairment. However, based on the results obtained in patients with severe renal impairment, a clinically significant increase in ritlecitinib exposure is not expected in these patients. The eGFR and classification of renal function status of participants was done using the Modification of Diet in Renal Disease (MDRD) formula.

Based on the above considerations, no dose adjustment is required in patients with mild, moderate or severe renal impairment. Ritlecitinib has not been studied in patients with ESRD or in renal transplant recipients (see section 4.2).

Hepatic impairment

Patients with moderate (Child Pugh B) hepatic impairment had an 18.5% increase in ritlecitinib AUC₂₄ compared to participants with normal hepatic function. Ritlecitinib was not studied in patients with mild (Child Pugh A) hepatic impairment. However, based on the results obtained in patients with moderate hepatic impairment, a clinically significant increase in ritlecitinib exposure is not expected in these patients. No dose adjustment is required in patients with mild or moderate hepatic impairment (see section 4.2). Ritlecitinib has not been studied in patients with severe (Child Pugh C) hepatic impairment (see section 4.3).

5.3 Preclinical safety data

General toxicity

Decreased lymphocyte counts and decreased lymphoid cellularity of organs and tissues of the immune and haematolymphopoietic systems were observed in nonclinical toxicity studies and were attributed to the pharmacological properties (JAK3/TEC inhibition) of ritlecitinib.

Chronic administration of ritlecitinib to Beagle dogs led to the occurrence of axonal dystrophy at systemic exposures of at least 7.4-times the expected exposure in patients treated with 50 mg per day (based on unbound AUC24). Axonal dystrophy is presumably related to binding to off-target neuronal proteins. It is not known if axonal dystrophy occurred in dogs at lower systemic exposures. At a systemic exposure that was 33-times above the expected exposure in patients treated with 50 mg per day (based on unbound AUC24), axonal dystrophy was associated with neurological hearing loss. While these findings proved to reverse after dosing cessation of ritlecitinib in dogs, a risk to patients at a chronic dosing regimen cannot be fully excluded (see section 4.4).

Genotoxicity

Ritlecitinib was not mutagenic in the bacterial mutagenicity assay (Ames assay). Ritlecitinib is not aneugenic or clastogenic at exposures equal to 130 times the MRHD on an unbound AUC basis based on the results of the *in vivo* rat bone marrow micronucleus assay.

Carcinogenicity

No evidence of tumorigenicity was observed in the 6-month Tg.ras H2 mice administered ritlecitinib at exposures equal to 11 times the MRHD on an unbound AUC basis. In a 2-year rat carcinogenicity study, a higher incidence of benign thymomas in female rats and benign thyroid follicular adenomas in male rats was noted following ritlecitinib administration at exposures equal to 29 times the MRHD on an unbound AUC basis. At this ritlecitinib exposure, a higher incidence of malignant thymomas in female rats cannot be excluded. No ritlecitinib-related thymomas or thyroid follicular adenomas were observed at exposures equal to 6.3 times the MRHD on an unbound AUC basis.

Reproductive and developmental toxicity

Ritlecitinib had no effects on female rat fertility at exposures equal to 55 times the MRHD on an unbound AUC basis. Effects on male rat fertility were noted (higher preimplantation loss resulting in lower number of implantation sites and corresponding lower litter size in naïve females mated with ritlecitinib dosed males) at exposure equal to 55 times the MRHD on an unbound AUC basis. No effects on male fertility were noted at exposures equal to 14 times the MRHD on an unbound AUC basis. No effects on spermatogenesis (sperm counts, sperm production rate, motility, and morphology) were noted at any dose in the rat fertility study.

In an embryo-foetal development study in pregnant rats, oral administration of ritlecitinib from gestation days 6 to 17 resulted in foetal skeletal malformations and variations and lower foetal body weights at exposures greater than or equal to 49 times the unbound AUC at the MRHD (see section 4.3). There were no effects on embryo-foetal development at exposures equal to 16 times the unbound AUC at the MRHD.

In an embryo-foetal development study in pregnant rabbits, oral administration of ritlecitinib from gestation days 7 to 19 resulted in lower mean foetal body weights and higher incidences of visceral malformations, skeletal malformations, and skeletal variations at exposures equal to 55 times the unbound AUC at the MRHD (see section 4.3). There were no effects on embryo-foetal development at exposures equal to 12 times the unbound AUC at the MRHD.

In a rat pre- and postnatal development study, oral administration of ritlecitinib from gestation day 6 through lactation day 20 resulted in developmental toxicity that included lower postnatal survival, lower offspring body weights, and secondary developmental delays at exposure equal to 41 times the unbound AUC at the MRHD (see section 4.3). Bred females in the F1 generation exhibited lower mean numbers of corpora lutea at exposures equal to 41 times the unbound AUC at the MRHD. There were no effects on pre- and postnatal development at exposures equal to 14 times the unbound AUC at the MRHD.

In a juvenile rat toxicity study, oral administration of ritlecitinib from postnatal day 10 to 60 (comparable to infant through adolescence human age) was not associated with effects on the nervous or skeletal systems.

Lactation

Following administration of ritlecitinib to lactating rats, concentrations of ritlecitinib in milk over time were higher than those in plasma, where the mean milk to plasma AUC ratio was determined to be 2.2 (see section 4.3).

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Hard capsule content

Cellulose microcrystalline Lactose monohydrate Crospovidone Glycerol dibehenate

Hard capsule shell

Hypromellose (E464) Titanium dioxide (E171) Yellow iron oxide (E172) Brilliant Blue FCF (E133)

Printing ink

Shellac Propylene glycol Ammonia solution concentrated Black iron oxide (E172) Potassium hydroxide

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

30 months.

6.4 Special precautions for storage

This medicinal product does not require any special temperature storage conditions. Store in the original package in order to protect from light.

6.5 Nature and contents of container

High-density polyethylene (HDPE) bottle with a silica gel desiccant and polypropylene closure containing 28 hard capsules.

OPA/Al/PVC/Al blisters containing 10 hard capsules. Each pack contains 30 or 90 hard capsules.

Not all pack sizes may be marketed.

6.6 Special precautions for disposal and other handling

Any unused medicinal product or waste material should be disposed of in accordance with local requirements.

7. MARKETING AUTHORISATION HOLDER

Pfizer Europe MA EEIG Boulevard de la Plaine 17 1050 Bruxelles Belgium

8. MARKETING AUTHORISATION NUMBER(S)

EU/1/23/1755/001 EU/1/23/1755/002 EU/1/23/1755/003

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

Date of first authorisation:

10. DATE OF REVISION OF THE TEXT

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

Pfizer Manufacturing Deutschland GmbH Betriebsstätte Freiburg Mooswaldallee 1 79090 Freiburg Germany

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 (PSURs)

The requirements for submission of PSURs for this medicinal product are set out in the list of Union reference dates (EURD list) provided for under Article 107c(7) of Directive 2001/83/EC and any subsequent updates published on the European medicines web-portal.

The marketing authorisation holder (MAH) shall submit the first PSUR for this product within 6 months following authorisation.

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

• Risk management plan (RMP)

The marketing authorisation holder (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.

Additional risk minimisation measures

Prior to the launch of ritlecitinib in each Member State, the MAH must agree about the content and format of the educational programme, including communication media, distribution modalities, and any other aspects of the programme, with the National Competent Authority.

The educational programme is aimed at increasing awareness about the safety concerns of the product, specifically in regard to infections (including herpes zoster and serious infections and opportunistic infections), thromboembolic events including deep vein thrombosis, pulmonary embolism and arterial thrombosis, MACE, malignancy, neurotoxicity and embryo-foetal toxicity following exposure *in utero*.

The MAH shall ensure that in each Member State where ritlecitinib is marketed, all healthcare professionals and patients/carers who are expected to prescribe, dispense or use ritlecitinib have access to/are provided with the following educational package:

The physician educational material should contain:

- The Summary of Product Characteristics
- Package leaflet
- Healthcare Professional Guide
- Patient Card (PC)

The healthcare professional guide shall contain the following key elements:

- Language for healthcare providers (HCPs) to inform patients of the importance of the PC.
- Potential risk of infections (including herpes zoster and serious infections or opportunistic infections)
 - O Describe that Litfulo must not be used in patients with an active, serious infection.
 - o Language on the risk of infections during treatment with Litfulo.
 - Language recommending that risk factors for infections should be considered when prescribing ritlecitinib including elderly age and diabetes.
 - Details on how to reduce the risk of infection with specific clinical measures (what laboratory parameters should be used to initiate Litfulo, screening for TB, and screening for viral hepatitis and temporary interruption of Litfulo if an infection is not responding to appropriate therapy until the infection is controlled).
 - o Language stating the use of live, attenuated vaccines should be avoided during or immediately prior to treatment along with examples of live, attenuated vaccines.
- Potential risk of thromboembolic events including deep vein thrombosis, pulmonary embolism and arterial thrombosis
 - Language describing that events of venous and arterial thromboembolism, including MACE, have been observed in studies in Litfulo.
 - Details of how to reduce the potential risk: Litfulo should be used with caution in patients with known risk factors for thromboembolism. In patients with a suspected thromboembolic event, discontinuation of Litfulo and prompt re-evaluation is recommended. The risks and benefits of treatment should be considered prior to initiating Litfulo therapy in patients.
- Potential risk of malignancy
 - Language describing that malignancies, including non-melanoma skin cancer, have been observed in studies with Litfulo.
 - Obetails of how to reduce the potential risk with specific clinical measures (that the risks and benefits of Litfulo treatment should be considered prior to initiating in patients with a known malignancy or when considering continuing Litfulo therapy in patients who develop a malignancy, and that periodic skin examination is recommended for patients who are at increased risk for skin cancer).
- Potential risk of neurotoxicity
 - Language describing that ritlecitinib-related axonal dystrophy has been observed in chronic Beagle dog toxicity studies at systemic exposures of at least 7.4-times the expected exposure in patients treated with 50 mg per day. At a systemic exposure that was 33-times above the expected exposure in patients treated with 50 mg per day, axonal dystrophy was associated with neurological hearing loss. While these findings proved to reverse after dosing cessation of ritlecitinib in dogs, a risk to patients at a chronic dosing regimen cannot be fully excluded. Available clinical data has not indicated an effect on neurological or audiological outcomes.
 - Details on how to reduce the risk Neurotoxicity, treatment with Litfulo should be discontinued in case unexplained neurological symptoms occur.
- Potential risk of embryo-foetal toxicity following exposure in utero
 - o Language describing there are no or limited data on the use of Litfulo in pregnant women.
 - Details on how to reduce the risk of exposure during pregnancy for women of childbearing potential based on the following: Litfulo is contraindicated during pregnancy,

women of childbearing potential should be advised to use effective contraception both during treatment and for 1 month following cessation of Litfulo, and to advise patients to inform their HCP immediately if they think they could be pregnant or if pregnancy is confirmed.

The patient information pack should contain:

- Package leaflet
- Patient card
- The patient card shall contain the following key messages:
 - o Language describing Litfulo (i.e. what it is and what it is used for).
 - o Contact details of the Litfulo prescriber.
 - o Language that the PC should be carried by the patient at any time and to share it with HCPs involved in their care (i.e., non- Litfulo prescribers, emergency room HCPs, etc.).
 - O Description of signs/symptoms of infections the patient needs to be aware of, so that they can seek attention from their HCP:
 - Language to advise patients and their HCPs about the risk of live vaccinations when given immediately before and during Litfulo therapy with examples of live vaccines.
 - o Reminder of the risk of cancer. Regarding skin cancer reminder to let their doctor know if they notice any new growth on the skin.
 - O Description of signs/symptoms of thromboembolic events including blood clots in the veins (deep vein thrombosis) or in the lungs (pulmonary embolism) and blood clots in an artery (arterial thrombosis), in the heart (heart attack), in the brain (stroke) or in the eye (profound vision loss in one eye) which the patient needs to be aware of, so that they can seek immediate attention from an HCP.
 - Language that treatment with Litfulo should be discontinued in case unexplained neurological symptoms occur.
 - o Language that there are no or limited data on the use of Litfulo in pregnant women.
 - Language describing on how to reduce the risk of exposure during pregnancy for women of childbearing potential based on the following:
 - Litfulo is contraindicated during pregnancy, women of childbearing potential should be advised to use effective contraception both during treatment and for 1 month following cessation of Litfulo, and to advise patients to inform their HCP immediately if they think they could be pregnant or if pregnancy is confirmed.
 - A reminder to use contraception, that Litfulo is contraindicated during pregnancy, and to notify their HCPs if they become pregnant while taking Litfulo.

ANNEX III LABELLING AND PACKAGE LEAFLET

A. LABELLING

PARTICULARS TO APPEAR ON THE OUTER PACKAGING
CARTON FOR BLISTER PACK FOR 50 MG
1. NAME OF THE MEDICINAL PRODUCT
Litfulo 50 mg hard capsules ritlecitinib
2. STATEMENT OF ACTIVE SUBSTANCE(S)
Each hard capsule contains ritlecitinib tosylate equivalent to 50 mg ritlecitinib.
3. LIST OF EXCIPIENTS
Contains lactose monohydrate. (see leaflet for further information).
4. PHARMACEUTICAL FORM AND CONTENTS
30 hard capsules 90 hard capsules
5. METHOD AND ROUTE(S) OF ADMINISTRATION
Oral use. Do not split, crush or chew. Read the package leaflet before 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
Store in the 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
Boule	Europe MA EEIG evard de la Plaine 17 Bruxelles um
12.	MARKETING AUTHORISATION NUMBERS
	/23/1755/002 30 hard capsules /23/1755/003 90 hard capsules
13.	BATCH NUMBER
Lot	
14.	GENERAL CLASSIFICATION FOR SUPPLY
15.	INSTRUCTIONS ON USE
16.	INFORMATION IN BRAILLE
Litful	o 50 mg
17.	UNIQUE IDENTIFIER – 2D BARCODE
2D ba	arcode carrying the unique identifier included.
18.	UNIQUE IDENTIFIER - HUMAN READABLE DATA
PC SN NN	

PARTICULARS TO APPEAR ON THE OUTER PACKAGING	
OTTLE LABEL FOR 50 MG	
1. NAME OF THE MEDICINAL PRODUCT	
Litfulo 50 mg hard capsules ritlecitinib	
2. STATEMENT OF ACTIVE SUBSTANCE(S)	
Each hard capsule contains ritlecitinib tosylate equivalent to 50 mg ritlecitinib.	
3. LIST OF EXCIPIENTS	
Contains lactose monohydrate. (see leaflet for further information).	
4. PHARMACEUTICAL FORM AND CONTENTS	
28 hard capsules	
5. METHOD AND ROUTE(S) OF ADMINISTRATION	
Oral use. Do not split, crush or chew. Read the package leaflet before 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	
Do not swallow the desiccant.	
8. EXPIRY DATE	
EXP	
9. SPECIAL STORAGE CONDITIONS	

Store in the 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
Boul	er Europe MA EEIG evard de la Plaine 17 Bruxelles ium
12.	MARKETING AUTHORISATION NUMBER(S)
EU/1	/23/1755/001 28 hard capsules
13.	BATCH NUMBER
Lot	
14.	GENERAL CLASSIFICATION FOR SUPPLY
15.	INSTRUCTIONS ON USE
16.	INFORMATION IN BRAILLE
Litfu	lo 50 mg
17.	UNIQUE IDENTIFIER – 2D BARCODE
2D b	arcode carrying the unique identifier included.
18.	UNIQUE IDENTIFIER - HUMAN READABLE DATA
PC SN NN	

MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS
BLISTERS FOR 50 MG CAPSULES
1. NAME OF THE MEDICINAL PRODUCT
Litfulo 50 mg hard capsules ritlecitinib
2. NAME OF THE MARKETING AUTHORISATION HOLDER
Pfizer Europe MA EEIG (as MA holder logo)
3. EXPIRY DATE
EXP
4. BATCH NUMBER
Lot
5. OTHER

B. PACKAGE LEAFLET

Package leaflet: Information for the patient

Litfulo 50 mg hard capsules

ritlecitinib

This medicine is subject to additional monitoring. This will allow quick identification of new safety information. You can help by reporting any side effects you may get. See the end of section 4 for how to report side effects.

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.

In addition to this leaflet, your doctor will give you a patient card, which contains important safety information that you need to be aware of. Keep this patient card with you.

What is in this leaflet

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

1. What Litfulo is and what it is used for

Litfulo contains the active substance ritlecitinib. It is used to treat severe alopecia areata in adults and adolescents 12 years of age and older. Alopecia areata is a disease where the body's own immune system attacks hair follicles, causing inflammation that leads to hair loss on the scalp, face and/or other parts of the body.

Litfulo works by reducing the activity of enzymes called JAK3 and TEC kinases, which are involved in inflammation at the hair follicle. This reduces the inflammation, leading to hair regrowth in patients with alopecia areata.

2. What you need to know before you take Litfulo

Do not take Litfulo

- if you are allergic to ritlecitinib or any of the other ingredients of this medicine (listed in section 6).
- if you have a serious infection ongoing, including tuberculosis.
- if you have severe liver problems.
- if you are pregnant or breast-feeding (see the "pregnancy, contraception, breast-feeding and fertility" section).

Warnings and precautions

Talk to your doctor or pharmacist before and during treatment with Litfulo if you:

- have an infection (possible signs may be fever, sweating, chills, muscle aches, cough, shortness of breath, blood in your phlegm, weight loss, diarrhoea, stomach pain, burning when you urinate, urinating more often than usual, feeling very tired). Litfulo can reduce your body's ability to fight infections and so worsen an existing infection or make it more likely for you to get a new infection.
- if you have diabetes or are older than 65 years of age, you may have an increased risk of getting infections.
- have, or have had, tuberculosis or have been in close contact with someone with tuberculosis, or if you reside or travel in regions where tuberculosis is very common. Your doctor will test you for tuberculosis before starting Litfulo and may retest you during treatment.
- have ever had a herpes infection (such as chickenpox or shingles), because Litfulo may allow it to come back. Tell your doctor if you get a painful skin rash with blisters as this can be a sign of shingles.
- have ever had hepatitis B or hepatitis C. Your doctor will test you for hepatitis before starting Litfulo and may retest you during treatment.
- have cancer or have had any cancer it is not clear if Litfulo increases the risk of cancer, and your doctor will discuss with you if treatment with this medicine is appropriate and whether check-ups including regular skin checks will be necessary during treatment.
- have had blood clots in the veins of your legs (deep vein thrombosis) or lungs (pulmonary embolism). Tell your doctor if you get a painful swollen leg, chest pain, or shortness of breath as these can be signs of blood clots in the veins.
- have had blood clots in an artery in the eye (retinal occlusion) or heart (heart attack). Tell your doctor if you experience acute changes to your eyesight (blurry vision, partial or complete loss of vision), chest pain, shortness of breath as these changes may be a sign of blood clots in the arteries.
- have recently had or plan to have a vaccination (immunisation) this is because certain vaccines (live vaccines) are not recommended while using Litfulo. Check with your doctor to see if your vaccinations are up to date and if you require additional vaccinations, including vaccination for shingles, before treatment with Litfulo.
- have unexplained symptoms caused by a problem with the nervous system while taking Litfulo. Your doctor will discuss with you if treatment should be discontinued.

Additional monitoring tests

Your doctor will carry out blood tests to check if you have low white blood cell count or low platelet count before and approximately 4 weeks after starting Litfulo treatment and may adjust your treatment if necessary.

Children

This medicine is not approved for use in children below the age of 12 years because the safety and benefits of Litfulo are not established in this age group.

Other medicines and Litfulo

Tell your doctor or pharmacist if you are taking, have recently taken or might take any other medicines.

In particular, tell your doctor or pharmacist before taking Litfulo if you are taking some of the medicines to treat:

- anxiety or sleep disorders (such as midazolam),
- heart rhythm problems (such as quinidine),
- gout (such as colchicine),
- rejection in organ transplantation (such as cyclosporine, everolimus, tacrolimus and sirolimus),
- migraine (such as dihydroergotamine and ergotamine),
- schizophrenia and chronic psychosis (such as pimozide),
- asthma (such as theophylline),

- muscle spasms (such as tizanidine),
- idiopathic pulmonary fibrosis (such as pirfenidone).

Litfulo may increase the amount of these medicines in your blood.

If any of the above apply to you or if you are not sure, talk to your doctor or pharmacist before taking Litfulo.

Pregnancy, contraception, breast-feeding and fertility

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.

Contraception in women

If you are a woman of childbearing potential, you should use an effective method of contraception during treatment with Litfulo, and for at least one month after your last treatment dose. Your doctor can advise you on suitable methods of contraception.

Pregnancy

Do not use Litfulo if you are pregnant, think you may be pregnant or are planning to have a baby. This medicine can harm the developing baby. Tell your doctor right away if you become pregnant or think you might have become pregnant during treatment.

Breast-feeding

Do not use Litfulo while breast-feeding as it is not known if this medicine passes into breast milk or if breast-fed babies are affected. You and your doctor should decide if you will breast-feed or use this medicine.

Fertility

It is unknown if Litfulo reduces fertility in women or men of childbearing potential.

Driving and using machines

Litfulo has no or limited effect on the ability to drive or use machines.

Litfulo contains lactose

If you have been told by your doctor that you have an intolerance to some sugars, contact your doctor before taking this medicine.

3. How to take Litfulo

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

The recommended dose is 50 mg once a day taken by mouth.

You should swallow your capsule whole with water. Do not open, crush or chew the capsule before swallowing as it may change how much medicine gets into your body.

You can take the capsule either with or without food.

If you take more Litfulo than you should

If you take more Litfulo than you should, contact your doctor. You may get some of the side effects described in section 4.

If you forget to take Litfulo

- If you miss a dose, take it as soon as you remember, unless your next dose is due in less than 8 hours.
- If there is less than 8 hours before your next dose, just skip the missed dose and take your next dose as usual when it is due.
- Do not take a double dose to make up for a forgotten capsule.

If you stop taking Litfulo

You should not stop taking Litfulo without discussing this with your doctor.

If you need to stop taking Litfulo for a short time (not more than 6 weeks), the risk of losing your scalp hair is low.

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 side effects

Talk to your doctor and get medical help straight away if you get any signs of:

- Shingles (herpes zoster), a painful skin rash with blisters with or without fever
- Hives (urticaria), an itching skin rash

Other side effects

Common (may affect up to 1 in 10 people)

- Infections of nose, throat or the windpipe
- Diarrhoea
- Dizziness
- Acne
- Rash (other than hives and shingles)
- Inflammation (swelling) of the hair follicles which may be itchy or painful (folliculitis)
- Increase in an enzyme called creatine phosphokinase, shown by blood test (blood creatine phosphokinase increased)

Uncommon (may affect up to 1 in 100 people)

- Low platelet count shown by blood test (platelet count decreased)
- Low white blood cell count shown by blood test (lymphocyte count decreased)
- Increase of liver enzymes in the blood (ALT and AST increased)

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 Litfulo

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, bottle, or blister after EXP. The expiry date refers to the last day of that month.

This medicine does not require any special temperature storage conditions. 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 throw away medicines you no longer use. These measures will help protect the environment.

6. Contents of the pack and other information

What Litfulo contains

- The active substance is ritlecitinib.

 Each hard capsule contains ritlecitinib tosylate equivalent to 50 mg ritlecitinib.
- The other ingredients are:

Hard capsule content: cellulose microcrystalline, lactose monohydrate, crospovidone), glycerol dibehenate (see section 2 "Litfulo contains lactose monohydrate").

Hard capsule shell: hypromellose (E464), titanium dioxide (E171), yellow iron oxide (E172), brilliant blue FCF (E133).

Printing ink: shellac, propylene glycol, ammonia solution concentrated, black iron oxide (E172), potassium hydroxide.

What Litfulo looks like and contents of the pack

Litfulo 50 mg opaque hard capsules have a yellow body and blue cap approximately 16 mm long and 6 mm wide of which the body is printed with "RCB 50" and the cap is printed with "Pfizer" in black.

The 50 mg hard capsules are provided in high-density polyethylene (HDPE) bottles with polypropylene closure containing 28 hard capsules or in OPA/Al/PVC/Al blisters containing 30 or 90 hard capsules. The bottle contains a silica gel desiccant used to keep the capsules dry. Do not swallow the silica gel desiccant.

Not all pack sizes may be marketed.

Marketing Authorisation Holder

Pfizer Europe MA EEIG Boulevard de la Plaine 17 1050 Bruxelles Belgium

Manufacturer

Pfizer Manufacturing Deutschland GmbH Betriebsstätte Freiburg Mooswaldallee 1 79090 Freiburg Germany For any information about this medicine, please contact the local representative of the Marketing Authorisation Holder:

België/Belgique/Belgien Luxembourg/Luxemburg

Pfizer NV/SA

Tél/Tel: +32 (0)2 554 62 11

България

Пфайзер Люксембург САРЛ, Клон България

Тел.: +359 2 970 4333

Česká republika

Pfizer, spol. s r.o. Tel: +420 283 004 111

Danmark

Pfizer ApS

Tlf: +45 44 20 11 00

Deutschland

PFIZER PHARMA GmbH

Tel: +49 (0)30 550055-51000

Eesti

Pfizer Luxembourg SARL Eesti filiaal

Tel: +372 666 7500

Ελλάδα

Pfizer Ελλάς Α.Ε. Τηλ: +30 210 6785800

España

Pfizer, S.L.

Tel: +34 91 490 99 00

France

Pfizer

Tél: +33 (0)1 58 07 34 40

Hrvatska

Pfizer Croatia d.o.o.

Tel: +385 1 3908 777

Ireland

Pfizer Healthcare Ireland

Tel: +1800 633 363 (toll free)

Tel: +44 (0)1304 616161

Ísland

Icepharma hf.

Sími: +354 540 8000

Lietuva

Pfizer Luxembourg SARL filialas Lietuvoje

Tel: +370 5 251 4000

Magyarország

Pfizer Kft.

Tel.: + 36 1 488 37 00

Malta

Vivian Corporation Ltd.

Tel: +356 21344610

Nederland

Pfizer by

Tel: +31 (0)800 63 34 636

Norge

Pfizer AS

Tlf: +47 67 52 61 00

Österreich

Pfizer Corporation Austria Ges.m.b.H.

Tel: +43 (0)1 521 15-0

Polska

Pfizer Polska Sp. z o.o.

Tel.: +48 22 335 61 00

Portugal

Laboratórios Pfizer, Lda.

Tel: +351 21 423 5500

România

Pfizer Romania S.R.L.

Tel: +40 (0) 21 207 28 00

Slovenija

Pfizer Luxembourg SARL

Pfizer, podružnica za svetovanje s področja

farmacevtske dejavnosti, Ljubljana

Tel: +386 (0)1 52 11 400

Slovenská republika

Pfizer Luxembourg SARL, organizačná zložka

Tel: + 421 2 3355 5500

Suomi/Finland

Pfizer Ov

Puh/Tel: +358 (0)9 430 040

Italia

Pfizer S.r.l.

Tel: +39 06 33 18 21

Κύπρος

Pfizer Ελλάς A.E. (Cyprus Branch)

Τηλ: +357 22817690

Latvija

Pfizer Luxembourg SARL filiāle Latvijā

Tel: + 371 670 35 775

This leaflet was last revised in .

Sverige Pfizer AB

Tel: +46 (0)8 550 520 00

United Kingdom (Northern Ireland)

Pfizer Limited

Tel: +44 (0) 1304 616161

Other sources of information

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