Olican® (fingolimod)
Healthcare Professional
Information

Olican® (fingolimod) Prescriber Patient Checklist

Important points to remember before, during and after treatment – Summary of Recommendations







Considerations in Olican® (fingolimod) Patient Selection

Olican® (fingolimod) is indicated as single disease modifying therapy in highly active relapsing remitting multiple sclerosis (MS) for the following groups of adult patients. While many patients are suitable for treatment, the following section highlights patients in whom Olican® (fingolimod) is contraindicated or not recommended.

Fingolimod causes transient heart rate reduction and may cause atrioventricular (AV) conduction delays following initiation of treatment. All patients should be monitored for a minimum of 6 hours on treatment initiation. Below is a brief overview of monitoring requirements.

Olican® (fingolimod) is generally recommended in MS patients who have had an inadequate response to, or are unable to tolerate, one or more therapies for multiple sclerosis.

Contraindications

Olican® (fingolimod) is contraindicated in patients with:

- hypersensitivity to fingolimod hydrochloride or to any of the excipients in the formulation of Olican® (fingolmod) or component of the container
- increased risk for opportunistic infections, including those who are immunocompromised due to treatment (e.g. antineoplastic, immunosuppressive or immunomodulating therapies, total lymphoid irradiation or bone marrow transplantation) or disease (e.g. immunodeficiency syndrome)
- severe active infections, including active chronic bacterial, fungal or viral infections (e.g. hepatitis, turberculosis)
- known active malignancies (except patients with basal cell carcinoma (BCC)
- in the previous 6 months, myocardial infarction (MI), unstable angina pectoris, stroke/transient ischaemic attack (TIA), decompensated heart failure (requiring inpatient treatment), or New York Heart Association (NYHA) class III/IV heart failure





The following patients should not be treated with Olican[®] (fingolimod)

- Those who are pregnant or breast feeding
- Those who are taking Class Ia or Class III antiarrhythmics
- Pediatric patients

Should not be used

Consider fingolimod only after performing risk/benefit analysis and consulting a cardiologist

Consult cardiologist regarding appropriate first-dose monitoring

History of bradyarrhythmia[†], QTc prolongation > 470 msec (females) or > 450 msec (males), or risk factors for QT prolongation, severe sleep apnoea, significant cardiovascular disease [‡], uncontrolled hypertension, cerebrovascular disease, history of recurrent syncope At least overnight extended monitoring is recommended

Consult cardiologist regarding possibility of switching to non-heart-rate-lowering drugs

Taking beta-blockers, heart-rate-lowering calcium channel blockers § or other substances that are known to lower the heart rate ||

If change inmedication is not possible, extend monitoring to at least overnight

- † Bradyarrhythmia includes the following: second-degree Mobitz type II or higher atrioventricular (AV) block, sick sinus syndrome, sinoatrial heart block, history of symptomatic bradycardia.
- ‡ Significant cardiovascular disease includes the following: ischaemic heart disease (including angina pectoris), history of myocardial infarction, congestive heart failure, history of cardiac arrest.
- Such as verapamil or diltiazem
- || Includes digoxin, cholinesterase inhibitors, or pilocarpine.





Patient's name: Date of birth:						
	Prior to initiating treatment					
	Ensure patients are not concomitantly taking Class Ia or Class III antiarrhythmic medicines					
	Conduct baseline electrocardiogram (ECG) and blood pressure measurement					
	Treatment with Olican® (fingolimod) should not be used in the following patients, unless anticipated benefits outweigh the potential risks					
	 Those with bradyarrhythmia¹, significant cardiovascular disease *, significant QT-interval prolongation, uncontrolled hypertension, cerebrovascular disease, severe untreated sleep apnoea, or a history of recurrent syncope Seek advice from a cardiologist regarding the most appropriate monitoring at treatment initiation; at least overnight extended monitoring is recommended] 					
	 Those receiving concurrent therapy with beta-blockers, heart-rate-lowering calcium channel blockers (such as verapamil or diltiazem), or other substances which may decrease heart rate (e.g., digoxin, cholinesterase inhibitors, or pilocarpine) Seek advice from a cardiologist regarding a switch to non-heart-rate-lowering medicinal products prior 					
	to initiation of treatment If heart-rate-lowering medication cannot be stopped, seek advice from a cardiologist regarding the most appropriate monitoring at treatment initiation; at least overnight extended monitoring is recommended appropriate monitoring at treatment initiation; at least overnight extended monitoring is recommended					





Patient's name: Date of birth:	
	Avoid co-administration of anti-neoplastic, immunosuppressive or immunomodulatory therapies due to the risk of additive immune system effects. For the same reason, a decision to use prolonged concomitant treatment with corticosteroids should be taken after careful consideration
	Obtain recent (within 6 months) transaminase, and bilirubin levels
	Obtain recent (within 6 months or after discontinuation of prior therapy) peripheral lymphocyte count (complete blood count (CBC). Treatment should not be initiated when lymphocyte counts are consistently below normal range.
	Confirm a negative pregnancy test result in women of childbearing potential
	Counsel on the need for effective contraception in women of childbearing age up to at least 2 months after completion or discontinuation of treatment, due to teratogenic risk to fetus
	Delay initiation of treatment in patients with severe active infection until resolved
	Check varicella zoster virus (VZV) antibody status in patients without a healthcare-professional-confirmed history of chickenpox or documentation of a full course of vaccination with varicella vaccine is recommended and treatment initiation should be delayed for 1 month to allow full effect of vaccination to occur
	Conduct an ophthalmologic evaluation in patients with history of uveitis or diabetes mellitus
	Conduct a dermatologic examination. The patient should be referred to a dermatologist if suspicious lesions, potentially indicative of basal cell carcinoma, are detected
	Provide patients with a Patient Reminder Card





[¶] Bradyarrhythmia includes the following: second-degree Mobitz type II or higher AV block, sick sinus syndrome, sinoatrial heart block, history of symptomatic bradycardia.

[#] Significant cardiovascular disease includes the following: ischaemic heart disease (including angina pectoris), history of myocardial infarction, congestive heart failure, history of cardiacarrest.

Patient's name: Date of birth:				
First-dose monitoring up to 6 hours post-dose and beyond, as necessary				
	Perform ECG and blood pressure 6 hours after the first dose			
	Monitor for signs and symptoms of bradyarrhythmia with hourly pulse and blood pressure measurements at least up to 6 hours post-dose. Initiate continuous monitoring as required.			
	Extend monitoring if: • Heart rate at 6 hours is < 45 bpm or is lowest post-dose value • Heart rate 6 at hours shows new onset second degree or higher AV block • If ECG at 6 hours shows QTc ≥ 500 msec. (overnight monitoring required)			
	Counsel patients that their ability to drive and use machines may be affected during and potentially after this period.			
During treatment				
	Conduct a full ophthalmologic evaluation in all patients at 3 to 4 months after starting treatment and in any patient complaining of visual disturbances >> Conduct periodic ophthalmologic evaluations in patients with history of uveitis or diabetes mellitus >> Counsel patients to report any visual disturbance during treatment			

>> Evaluate the fundus, including the macula, and discontinue treatment if macular oedema is confirmed





Patie	nt's name:	Date of birth:
	>> >> >>	Prompt antimicrobial treatment should be initiated if indicated Perform prompt diagnostic evaluation in patients with symptoms and signs consistent with cryptococcal meningitis, and initiate appropriate treatment if diagnosed (cryptococcal meningitis, sometimes fatal, 2-3 years after treatment ggestive of progressive multifocal leukoencephalopathy (PML). If PML is suspected, treatment with fingolimod should be suspended until PML has been excluded (cases of PML 2-3 years after treatment) If a patient develops a serious infection, treatment should be suspended Discontinue treatment in disseminated herpetic infections.
	thereafter. T	mphocyte count (CBC) should be monitored during treatment, at month 3 and at least yearly reatment should be interrupted if lymphocyte count is confirmed as <0.2x109/L. The approved 5 mg once daily when restarting Olican® (fingolimod) should be administered. Other dosing regimens gilant for clinical symptoms or MRI findings that may be su
	Check liver transaminases and serum bilirubin before starting treatment and at months 1, 3, 6, 9, and 12 and periodically thereafter, until 2 months after fingolimod discontinuation, or at any time there are signs or symptoms of hepatic dysfunction In the absence of clinical symptoms, if liver transaminases are: >> Greater than 3 times the upper limit of normal (ULN) but less than 5 times ULN without increase in serum bilirubin, more frequent monitoring including serum bilirubin and alkaline phosphatase (ALP)	
	,	should be instituted.





Patien	atient's name: Date of birth:		
	>> At least 5 times ULN or at least 3 times ULN associated with any increase in serum bilirubin, fingolimod should be discontinued. If serum levels return to normal, fingolimod may be restarted based on a careful benefit-risk assessment of the patient.		
	In the presence of clinical symptoms suggestive of hepatic dysfunction:		
	>> Liver enzymes and bilirubin should be checked promptly and fingolimod should be discontinued if significant liver injury is confirmed.		
	During treatment and for up to 2 months after discontinuation >> Vaccinations may be less effective >> Live attenuated vaccines may carry a risk of infection and should be avoided		
	Counsel patient to advise physician immediately if she becomes pregnant Pregnancy tests should be repeated at suitable intervals. Discontinue treatment if a patient becomes pregnant To help determine the effects of fingolimod exposure in pregnant women with MS, physicians are encouraged to report any pregnancy outcomes by calling the National Pharmacovigilance Centre (NPC) at +966-11-2038222, Ext:2317-2356-2340		
	Vigilance for basal cell carcinoma and other cutaneous neoplasms is recommended, with skin examination prior to treatment initiation and then yearly taking into consideration clinical judgment and referral to a dermatologist if suspicious lesions, potentially indicative of basal cell carcinoma or other cutaneous neoplasms, are detected. Caution patients against exposure to sunlight without protection. These patients should not receive concomitant phototherapy with UV-B-radiation or PUVA-photochemotherapy.		





Patient's name:			
	After treatment discontinuation		
	Repeat first-dose monitoring as for treatment initiation when treatment is interrupted for: >> One day or more during the first 2 weeks of treatment >> More than 7 days during weeks 3 and 4 of treatment >> More than 2 weeks after 1 month of treatment		
	Counsel patients to report signs and symptoms of infection for up to 2 months after discontinuation		
	Counsel patients that effective contraception is needed for 2 months after discontinuation		

Treatment initiation algorithm

All patients will need to be monitored for at least 6 hours during treatment initiation, as described in the algorithm below. In addition, for patients in whom fingolimod is not recommended, advice should be sought from a cardiologist regarding appropriate monitoring; at least overnight monitoring is recommended for this group.





Monitor for a minimum of 6 hours

- Perform ECG and BP measurement
- Monitor for a minimum of 6 hours for signs and symptoms of bradycardia, with hourly pulse and BP checks. If patient is symptomatic, continue monitoring until resolution
 - Continuous (real-time) ECG is recommended throughout the 6-hour period
- **¤** Perform ECG at 6 hours

Treatment initiation algorithm

Did the patient require pharmacologic intervention at any time during the monitoring period?



Monitor overnight.

First-dose monitoring should be repeated after the second dose of Olican® (fingolimod)



Did third-degree AV block occur at any time during the monitoring period?





At the end of the monitoring period, have any of the following criteria been met?

- **¤** HR < 45 bpm
- □ ECG shows new-onset second-degree or higher AV block or QTc interval ≥ 500 msec



Extent monitoring at least overnight, until the findings have resolved







Treatment initiation algorithm (cont.)



At the end of the monitoring period, is the HR the lowest since the first dose was administered?



Extent monitoring for at least 2 hours and until heart rate increases



First-dose monitoring is complete



The above first-dose monitoring procedure should also be followed at reinitiation of treatment if Olican® (fingolimod) therapy is discontinued for

- One day or longer within the first 2 weeks of treatment
- More than 7 days during weeks 3 and 4 of treatment
- More than 2 weeks after the first month of treatment.

BP= blood pressure; ECG=electrocardiogram; HR= heart rate; QTc= heart-rate-corrected QT interval





Olican® (fingolimod) Healthcare Professional Information

Summary of SPC to be added.





This document is approved by The Executive Directorate of Pharmacovigilance, at SFDA.

For extra copies you can contact Pharmascience at

Telephone: +966 11 293 1841

Reporting adverse events:

Healthcare professionals should report any suspected adverse reactions associated with the use of Olican® (fingolimod). Therefore, if you receive or observe any adverse reaction you can reach the following contacts:

Saudi Food and Drug Authority, **National Pharmacovigilance Center**

Unified Contact Center: 19999 Email: npc.drug@sfda.gov.sa

Or by online: https://ade.sfda.gov.sa

Pharmacovigilance Department, Pharmascience Inc.:

Tel: +966 11 293 1841 Fax: +966 11 293 1353

Email: pms.sa@pharmascience.com

Version 2: February 2021





