



MHRA
10 South Colonnade
Canary Wharf
London
E14 4PU
United Kingdom

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### **Decision Cover Letter**

## Decision of the licensing authority to:

accept change(s) to the agreed paediatric investigation plan and to the deferral MHRA-100195-PIP01-21-M01  $\,$ 

## **Scope of the Application**

**Active Substance(s)** 

**BALOXAVIR MARBOXIL** 

Condition(s)

Treatment of Influenza, Prevention of Influenza

## **Pharmaceutical Form(s)**

Film-coated tablet, Granules for oral suspension

#### **Route(s) of Administration**

Oral use

## Name / Corporate name of the PIP applicant

**Roche Products Limited** 

#### **Basis for the Decision**

Pursuant to the Human Medicines Regulations 2012, Roche Products Limited submitted to the licensing authority on 05/11/2021 07:56 GMT an application for a Modification

The procedure started on 18/02/2022 10:02 GMT

1. The licensing authority, having assessed the application in accordance with the Human Medicines Regulations 2012, decides, as set out in the appended summary report:

to accept change(s) to the agreed paediatric investigation plan and to the deferral

2. The measures and timelines of the paediatric investigation plan are set out in the Annex I.

This decision is forwarded to the applicant, together with its annex and appendix.





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## **Final Decision Letter**

MHRA-100195-PIP01-21-M01

Of 24/02/2022 15:46 GMT

On the adopted decision for BALOXAVIR MARBOXIL (MHRA-100195-PIP01-21-M01) in accordance with the Human Medicines Regulations 2012.

The licensing authority, in accordance with the Human Medicines Regulations 2012, has adopted this decision:

Agreement on modification of a paediatric investigation plan (including modification of a waiver or deferral included in that paediatric investigation plan)

This decision applies to a Modification for BALOXAVIR MARBOXIL, Film-coated tablet, Granules for oral suspension , Oral use .

This decision is addressed to Roche Products Limited, 6 Falcon Way, Shire Park, Welwyn Garden City, United Kingdom, AL7 1TW

#### ANNEX I

- 1. Waiver
- 1.1 Condition:

Not applicable to Treatment of Influenza or Prevention of influenza

#### 2. Paediatric Investigation Plan:

#### 2.1 Condition(s):

Treatment of Influenza infection; Prevention of influenza

#### 2.2 Indication(s) targeted by the PIP:

Treatment of Influenza infection; Prevention of influenza

# 2.3 Subset(s) of the paediatric population concerned by the paediatric development:

The paediatric population from birth to less than 18 years of age

# **2.4 Pharmaceutical Form(s):**

Film-coated tablet; Granules for oral suspension

# 2.5 Studies:

Study Type	Number of Studies	Study Description		
Quality Measures	0	Not applicable		
Non-Clinical Studies	0	Not applicable		
Clinical Studies	4	(For Treatment of Influenza) Study 1		
		(CP40559) Multicentre, single-arm,		
		open label study to assess safety,		
		pharmacokinetics, and efficacy		
		of baloxavir marboxil in infants		
		from birth to less than 1 year of		
		age with influenza-like symptoms.		
		Study 2 (CP40563) Multicentre,		
		randomised, double-blind, active-		
		controlled study to assess the safety,		
		pharmacokinetics, and efficacy of		
		baloxavir marboxil compared to		
		oseltamivir in children from 1 year		
		to less than 12 years of age with		
		influenza-like symptoms. Study 3		
		(CP40617) Randomised, double-		
		blind, placebo-controlled, multicentre		
		study to assess efficacy, safety,		
		and pharmacokinetics of baloxavir		
		marboxil in combination with a		
		standard of care neuraminidase		
		inhibitor (SOC NAI), compared with		
		a matching placebo in combination		
		with a SOC NAI in adolescents who		
		require hospitalisation for severe		
		influenza, or who contract severe		
		influenza during hospitalisation		
		(and adults). Study 4 was removed		
		during procedure EMEA-002440-		
		PIP01-18-M01(For Prevention of		
		Influenza)Study 13 Randomised,		
		double-blind, multicentre,		
		parallel-group, placebo-controlled		
		comparative study to evaluate		
		the efficacy and safety of a single		
		oral dose of baloxavir marboxil in		
		the prevention of influenza virus		
		infection in paediatric subjects from		

	birth to less than 18 years of age who are household members of influenza-
Extrapolation, Modeling & Simulation Studies	infected index patients.  (For Treatment of Influenza) Study 5 (M-S Study 1) Population PK model in otherwise healthy and high risk adult and paediatric subjects, evaluating relevant demographic covariates that may influence systemic drug exposure. Study 6 was removed during procedure EMEA-002440-PIP01-18-M01 Study 7 (MS Study 3) Modelling and simulation study in Otherwise Healthy and High risk patients evaluating exposure-response using viral kinetic modelling and primary efficacy endpoints. Study 8 was removed during procedure EMEA-002440-PIP01-18-M01 Study 9 was removed during procedure EMEA-002440-PIP01-18-M01 Study 10 (Extrapolation study 1) Partial extrapolation of PK, PD (virology, virus titres) and efficacy (primary endpoint, time to symptom alleviation) in otherwise healthy paediatric patients from birth to less than 12 years of age. Study 11 (Extrapolation study 2) Complete extrapolation of PD and efficacy (primary endpoint, time to symptom alleviation) in High Risk paediatric patients from birth to less than 12 years of age. Study 12 was removed during procedure EMEA-002440-
	PIPUI-10-WIUI
Other Studies	PIP01-18-M01 Not applicable

# 3. Follow-up, completion and deferral of a PIP:

Concerns on potential long term safety and	No
efficacy issues in relation to paediatric use:	
Date of completion of the paediatric	31/05/2024
investigation plan:	
Deferral of one or more studies contained in	Yes
the paediatric investigation plan:	