

Medicines & Healthcare products Regulatory Agency

> 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-100087-PIP01-21-M02

Scope of the Application

Active Substance(s)

LACOSAMIDE

Condition(s)

Treatment of generalised epilepsy and epileptic syndromes.

Pharmaceutical Form(s)

Film-coated tablet; Syrup; Solution for infusion

Route(s) of Administration

Oral use, Intravenous use

Name / Corporate name of the PIP applicant

UCB Pharma Limited

Basis for the Decision

Pursuant to the Human Medicines Regulations 2012, UCB Pharma Limited submitted to the licensing authority on 21/02/2022 17:37 GMT an application for a Modification

The procedure started on 20/06/2022 10:25 BST

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-100087-PIP01-21-M02

Of 03/08/2022 17:06 BST

On the adopted decision for LACOSAMIDE (MHRA-100087-PIP01-21-M02) 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 LACOSAMIDE, Film-coated tablet; Syrup; Solution for infusion, Oral use, Intravenous use.

This decision is addressed to UCB Pharma Limited, 208 Bath Road, , Slough, Berkshire, United Kingdom, SL1 3WE

ANNEX I

1. Waiver

1.1 Condition:

Not applicable

2. Paediatric Investigation Plan:

2.1 Condition(s):

Treatment of generalised epilepsy and epileptic syndromes

2.2 Indication(s) targeted by the PIP:

Treatment of generalised epilepsy and epileptic syndromes

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; Syrup; Solution for infusion

2.5 Studies:

Study Type	Number of Studies	Study Description
Quality Measures	3	Measure 1 Confirmation of the
		age-appropriateness of the current
		commercial lacosamide film-coated
		tablets (for oral use) for the target
		population. Measure 2 Confirmation
		of the age-appropriateness of the
		current commercial lacosamide
		solution for infusion (for intravenous
		use) for the target population.
		Measure 3 Confirmation of the
		age-appropriateness of the current
		commercial lacosamide syrup (for
		oral use) for the target population.
Non-Clinical Studies	0	Not applicable
Clinical Studies	6	Study 12 Open-label, multicentre
		study to investigate the
		pharmacokinetics (PK) of lacosamide
		(commercially available tablet or
		oral solution) as therapy in children
		(aged from 1 month to less than 18
		years) who are prescribed lacosamide
		for epilepsy (SP1047). Study 14
		Exploratory, open-label, study in
		paediatric subjects from 1 month
		to less than 18 years for safety
		and tolerability and preliminary
		efficacy for adjunctive lacosamide
		treatment of epilepsy syndromes
		associated with generalised seizures
		excluding primary generalised tonic
		clonic seizures with Idiopathic
		Generalised Epilepsy and excluding
		typical absence (Type IIA1) or
		atypical absence (Type IIA2)
		seizures when occurring exclusively

		from other seizure types (SP0966).
		Study 15 Open-label, multi-centre, parallel-group, non-inferiority
		efficacy, safety, tolerability and PK
		study for adjunctive lacosamide
		treatment in neonates with repeated
		electroencephalographic neonatal
		seizures (SP0968). Study 17: deleted
		during procedure EMEA-000402-
		PIP03-17-M03. Study 18 Open label, long term safety, tolerability
		and pharmacokinetic study in
		children from 1 month to less than
		18 years with epilepsy; extension
		study for subjects from other LCM
		studies including SP847 and SP0966
		(SP848). Study 20 Double-blind,
		randomised, placebo-controlled,
		parallel group, multi-centre study to evaluate efficacy and safety of
		lacosamide as adjunctive treatment
		for uncontrolled primary generalised
		tonic-clonic (PGTC) seizures in
		subjects aged 4 years and above
		with idiopathic generalised epilepsy
		(IGE) (SP0982). Study 21 Open-
		label, multi-centre, extension study
		to evaluate safety and tolerability of lacosamide as adjunctive treatment
		for idiopathic generalised epilepsy
		(IGE) with uncontrolled primary
		generalized tonic-clonic (PGTC)
		seizures in subjects aged 4 years and
		above with idiopathic generalised
Extrapolation Modeling &	5	epilepsy (IGE) (EP0012).
Extrapolation, Modeling & Simulation Studies	5	Study 4 PBPK prediction of oral lacosamide pharmacokinetics and
Simulation Studies		dose adaptations in children from
		birth to less than 18 years (CL0096).
		Study 5 Population pharmacokinetics
		of lacosamide in children with
		partial onset seizures aged from 1
		month to less than 18 years, based in data from studies SP847 and
		SP1047. Study 6 Physiologically
		based pharmacokinetic (PBPK)
		prediction of intravenous lacosamide
		pharmacokinetics and dose
		adaptations in neonates (aged from
		birth to 28 days). Study 7 Predictive
		population pharmacokinetics of intravenous lacosamide in
		children from birth to less than 18
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		years. Study 8 Final retrospective population pharmacokinetics model of lacosamide in children from birth to less than 18 years, combining all available data at the end of the program.
Other Studies	0	Not applicable
Other Measures	0	Not applicable

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

Concerns on potential long term safety and	Yes
efficacy issues in relation to paediatric use:	
Date of completion of the paediatric	31/12/2024
investigation plan:	
Deferral of one or more studies contained in	Yes
the paediatric investigation plan:	