

**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-100558-PIP01-22-M02

### **Scope of the Application**

#### **Active Substance(s)**

milvexian

#### **Condition(s)**

Prevention of thromboembolism in patients with cardiovascular disease

#### **Pharmaceutical Form(s)**

Film coated tablets; Age appropriate oral dosage form

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

ORAL USE

#### **Name / Corporate name of the PIP applicant**

Janssen-Cilag Limited

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

Pursuant to the Human Medicines Regulations 2012, Janssen-Cilag Limited submitted to the licensing authority on 28/11/2025 13:21 GMT an application for a Modification

The procedure started on 02/12/2025 14:41 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-100558-PIP01-22-M02

Of 02/04/2026 13:43 BST

On the adopted decision for milvexian (MHRA-100558-PIP01-22-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 milvexian, Film coated tablets; Age appropriate oral dosage form , ORAL USE .

This decision is addressed to Janssen-Cilag Limited, 50-100 Holmers Farm Way Buckinghamshire, High Wycombe, UNITED KINGDOM, HP12 4EG

## ANNEX I

### 1. Waiver

#### 1.1 Condition:

Prevention of thromboembolism in patients with cardiovascular disease. The waiver applies / applied to: Paediatric Subset(s): The paediatric population from birth to less than 28 days of age  
Pharmaceutical form(s): Film-coated tablet Age appropriate oral solid dosage form Route(s) of administration: ORAL USE Reason for granting waiver: on the grounds that the specific medicinal product does not represent a significant therapeutic benefit over existing treatments

### 2. Paediatric Investigation Plan:

#### 2.1 Condition(s):

Prevention of thromboembolism in patients with cardiovascular disease

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

Primary prevention of thromboembolic events in paediatric patients from 28 days to less than 18 years of age with congenital heart disease

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

The paediatric population from 28 days to less than 18 years of age

## 2.4 Pharmaceutical Form(s):

Film-coated tablet Age appropriate oral solid dosage form

## 2.5 Studies:

| Study Type           | Number of Studies | Study Description                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          |
|----------------------|-------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Quality Measures     | 1                 | Study 1 Development of an age-appropriate oral solid formulation.                                                                                                                                                                                                                                                                                                                                                                                                                                          |
| Non-Clinical Studies | 3                 | Study 2 In vitro assessment of coagulation assays in paediatric plasma samples spiked with milvexian. Study 3 Reproductive toxicity animal study to assess potential effects of milvexian on pregnant/lactating rabbits and postnatal development of the offspring. Study 7 Added during procedure MHRA-100558-PIP01-22-M02 Juvenile toxicity study in rabbits to assess the potential toxicity of milvexian when administered orally.                                                                     |
| Clinical Studies     | 2                 | Study 4 Open-label, single dose trial to evaluate pharmacokinetics, safety, tolerability, acceptability, and palatability of milvexian in children from 28 days to less than 18 years of age at risk of thromboembolic events. Study 5 Open label, randomised, multiple dose trial to evaluate pharmacokinetics, safety, and efficacy of milvexian compared to best standard of care in children from 28 days to less than 18 years of age with congenital heart disease at risk of thromboembolic events. |

|                                                         |   |                                                                                                                                                                                                                                                                                                                                                                                                                  |
|---------------------------------------------------------|---|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <b>Extrapolation, Modeling &amp; Simulation Studies</b> | 2 | Study 6 Modelling and simulation study to inform the dose and treatment regimen in studies 4 and 5, to achieve comparable drug exposure between adult and paediatric populations and to characterise PK/PD relationships. Extrapolation Plan Studies 4, 5, 6 are part of an extrapolation plan covering the paediatric population from 28 days to less than 18 years of age, as agreed by the Regulatory Agency. |
| <b>Other Studies</b>                                    | 0 | Not applicable.                                                                                                                                                                                                                                                                                                                                                                                                  |
| <b>Other Measures</b>                                   | 0 | Not applicable.                                                                                                                                                                                                                                                                                                                                                                                                  |

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

|                                                                                                  |            |
|--------------------------------------------------------------------------------------------------|------------|
| <b>Concerns on potential long term safety and efficacy issues in relation to paediatric use:</b> | No         |
| <b>Date of completion of the paediatric investigation plan:</b>                                  | 31/12/2033 |
| <b>Deferral of one or more studies contained in the paediatric investigation plan:</b>           | Yes        |