

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-100093-PIP01-21-M02

## **Scope of the Application**

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

UPADACITINIB

#### Condition(s)

Treatment of chronic idiopathic arthritis (including rheumatoid arthritis, psoriatic arthritis, spondyloarthritis and juvenile idiopathic arthritis)

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

Prolonged release tablet; Age-appropriate oral liquid dosage form; Age-appropriate oral solid dosage form

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

Oral use

### Name / Corporate name of the PIP applicant

Abbvie Ltd

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

Pursuant to the Human Medicines Regulations 2012, Abbvie Ltd submitted to the licensing authority on 20/12/2021 22:15 GMT an application for a Modification

The procedure started on 31/08/2022 08:31 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-100093-PIP01-21-M02

Of 08/11/2022 10:47 GMT

On the adopted decision for UPADACITINIB (MHRA-100093-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 UPADACITINIB, Prolonged release tablet; Age-appropriate oral liquid dosage form; Age-appropriate oral solid dosage form, Oral use.

This decision is addressed to Abbvie Ltd, AbbVie House, Vanwall Business Park, Vanwall Road, Maidenhead, United Kingdom, SL6 4UB

# ANNEX I

1. Waiver

### **1.1 Condition:**

Treatment of chronic idiopathic arthritis (including rheumatoid arthritis, psoriatic arthritis, spondyloarthritis and juvenile idiopathic arthritis) The waiver applies / applied to: Paediatric Subset(s): The paediatric population from birth to less than 1 year of age Pharmaceutical form(s): Age-appropriate oral liquid dosage form; Age-appropriate oral solid dosage form; Prolonged release tablet Route(s) of administration: Oral use Reason for granting waiver: on the grounds that the disease or condition for which the specific medicinal product is intended does not occur in the specified paediatric subset(s).

### 2. Paediatric Investigation Plan:

### **2.1 Condition(s):**

Treatment of chronic idiopathic arthritis (including rheumatoid arthritis, psoriatic arthritis, spondyloarthritis and juvenile idiopathic arthritis)

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

Treatment of Juvenile Idiopathic Arthritis

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

The paediatric population from 1 year to less than 18 years of age

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

Age-appropriate oral liquid dosage form; Age-appropriate oral solid dosage form; Prolonged release tablet

### 2.5 Studies:

| Study Type                | Number of Studies | Study Description                      |
|---------------------------|-------------------|--|
| Quality Measures          | 1                 | Study 1 Development of age-            |
|                           |                   | appropriate oral solid dosage          |
|                           |                   | form (dispersible tablet or multi-     |
|                           |                   | particulate granules) or age-          |
|                           |                   | appropriate oral liquid dosage form.   |
| Non-Clinical Studies      | 2                 | Study 2 Dose range-finding juvenile    |
|                           |                   | toxicity study. Study 3 Definitive     |
|                           |                   | juvenile toxicity study to evaluate    |
|                           |                   | toxicity and impact of upadacitinib    |
|                           |                   | on neonatal/juvenile development.      |
| Clinical Studies          | 2                 | Study 4 Open-label, multiple           |
|                           |                   | dose study to evaluate the             |
|                           |                   | pharmacokinetics, safety, and          |
|                           |                   | tolerability and to confirm the dosing |
|                           |                   | regimen of upadacitinib in children    |
|                           |                   | with active polyarticular course JIA.  |
|                           |                   | Study 5 Deleted during procedure       |
|                           |                   | MHRA-100093-PIP01-21-M02.              |
|                           |                   | Study 6 Randomised, placebo-           |
|                           |                   | controlled, double-blind withdrawal    |
|                           |                   | study to evaluate the safety and       |
|                           |                   | efficacy of multiple doses of          |
|                           |                   | upadacitinib in children with active   |
|                           |                   | systemic JIA.                          |
| Extrapolation, Modeling & | 1                 | Study 7 Population pharmacokinetic     |
| Simulation Studies        |                   | two compartment model that             |
|                           |                   | characterises the pharmacokinetic      |
|                           |                   | parameters, the inter- and intra-      |

|                |   | subject variability, and relationship<br>between pharmacokinetic parameters<br>and the relevant covariates. |
|----------------|---|---|
| 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<br>efficacy issues in relation to paediatric use: | Yes        |
|--|------------|
| Date of completion of the paediatric investigation plan:                                     | 31/08/2027 |
| Deferral of one or more studies contained in the paediatric investigation plan:              | Yes        |