

To:

Head of Paediatric Medicines  
European Medicines Agency

**Notification of discontinuation of a paediatric development which is covered by an agreed PIP Decision**

Actives substances(s): Fusion protein composed of the first 2 immunoglobulin-like domains of the human roundabout guidance receptor 2 fused to a human IgG1 crystallised fragment (PF-06730512)

Invented name: n/a

Latest Decision number(s): 1) P/0431/2022

Corresponding PIP number(s): 1) EMEA-003157-PIP01-21

Date of initial marketing authorisation granted: n/a

Date of authorisation of new indication, pharmaceutical form or route of administration: n/a

Please note that development of the medicinal product above in the following **condition(s)/indication(s)**:

Treatment of focal segmental glomerulosclerosis

- has been discontinued
- has been suspended/put on long-term hold (with possible re-start at a later time)
- for the following reason(s): (tick all that apply)
- (possible) lack of efficacy in adults
- (possible) lack of efficacy in children
- (possible) unsatisfactory safety profile in adults
- (possible) unsatisfactory safety profile in children
- commercial reasons (please specify: )
- manufacturing / quality problems
- other regulatory action (please specify: ) (e.g. suspension, revocation of M.A.)
- other reason (please specify: )

Please add a brief description (max 2000 characters) of the reason(s) for the discontinuation / suspension:

Following the 3rd planned interim analysis of Study C0221002 (a PHASE 2, 24-WEEK, ADAPTIVE, OPEN LABEL, SEQUENTIAL COHORT TRIAL TO EVALUATE THE EFFICACY, SAFETY, TOLERABILITY AND PHARMACOKINETICS OF PF-06730512 FOLLOWING MULTIPLE DOSES IN ADULT SUBJECTS WITH FOCAL SEGMENTAL GLOMERULOSCLEROSIS [FSGS]), Pfizer has decided to terminate the study and

the PF-06730512 development program due to lack of efficacy at both tested doses in study C0221002. Please note that this decision is not related to a safety concern, as no safety signals have been associated with PF-06730512 to date.

Please note that if the PIP has been submitted as part of a marketing authorisation application in order to comply with the requirements of Article 7 of the Paediatric Regulation (as a condition of the validation of the respective application) and a marketing authorisation was granted based on this application, then there is a legal obligation to complete that PIP. The same applies if there has been a successful post-authorisation application, where the PIP was included in order to comply with the requirements of Article 8 of the Paediatric Regulation.

Please confirm if any of the above applies to the PIP in question:

Yes  No

If yes, it means that based on the Marketing Authorisation obtained at the end of that initial procedure or the successful post-authorisation application, as applicable, you are obliged to complete that PIP. That obligation cannot be cancelled by a unilateral decision, including by withdrawing the MA. Such PIP must be completed, unless it is modified in agreement with the PDCO by removing all outstanding PIP measures or granting a full product-specific waiver instead (upon relevant circumstances in accordance with the Paediatric Regulation). Non-completion of a binding PIP establishes noncompliance with the requirements of the Paediatric Regulation, which the European Medicines Agency has an obligation to report to the European Commission.

Name and signature of the PIP contact point: Signature on file

Date: 16 February 2024

Contact for inquiries from interested parties:

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