## Notification of discontinuation of a paediatric development which is covered by an agreed PIP Decision<sup>i</sup>

Actives substances(s): efavale	Actives substances(s): efavaleukin alfa		
Latest Decision number(s):	1) P/0432/2022		
Corresponding PIP number(s):	1) EMEA-003156-PIP01-21		
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 abo	ove applies:		
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.			
Please note that development condition(s)/indication(s):	of the medicinal product above in the following		
Treatment of Systemic Lupus Erythematosus (SLE)			
$oxed{\boxtimes}$ has been discontinued			
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 sat	fety profile in children		
☐ commercial reasons (please	specify: )		
manufacturing / quality pro	blems		
$\hfill \square$ other regulatory action	(please specify: )		
☑ other reason	(please specify: please see below)		

Please add a brief description (max 2000 char	racters) of the reason(s) for the discontinuation:
Amgen wishes to notify the Agency to discont systemic lupus erythematosus. This is due to a	inue the development of efavaleukin alfa for treatment of a sponsor decision and not safety related.
Date:	20 November 2024
Contact for inquiries from interested parties:	Amgen Europe B.V.
Telephone:	+ 44 (0)1223 420305
Email:	medinfointernational@amgen.com

This form will be published to the corresponding decision available on the website of the European Medicines Agency.