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): benraliz	zumab		
Invented name: Fasenra	ì		
Latest Decision number(s): 1) P/0038/2022			
Corresponding PIP number(s):	1) EMEA-001214-PIP	07-21	
Date of initial marketing author	risation granted: 8 Jan	uary 2018	
Date of authorisation of new indication, pharmaceutical form or route of administration: n/a			
Please note that development condition(s)/indication(s):	of the medicinal produ	ct above in the following	
Eosinophilic gastritis/gastroent	eritis		
$oxed{\boxtimes}$ 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 saf	ety profile in adults		
(possible) unsatisfactory saf	ety profile in children		
commercial reasons (please	specify: )		
manufacturing / quality prol	olems		
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 /	

There is a growing body of evidence of eosinophil independent mechanisms in the pathology of eosinophilic upper gastrointestinal (GI) disease, including eosinophilic esophagitis and EG/EGE. The

suspension:

emerging scientific evidence that had become available since the HUDSON GI Phase III trial was initiated suggested the study was unlikely to result in consistent clinically meaningful benefits for patients in the trial. Therefore, AstraZeneca made the decision to stop enrolment in the study. The decision to terminate the study was not related to safety concerns.

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.

requirements of Article 8 of the Paediatric Reg	ulation.	
Please confirm if any of the above applies to the	ne 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:	30 June 2023	
Contact for inquiries from interested parties:	AstraZeneca	
Telephone:	+46 8 553 244 00	
Email:	paediatrics@astrazeneca.com	