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): Recombinant human monoclonal antibody to GM-CSF (Otilimab)		
Invented name: not applicable		
Latest Decision number(s): 1) P/0266/2020		
Corresponding PIP number(s): 1) EMEA-001882-PIP02-16-M02		
Date of initial marketing authorisation granted: not applicable		
Date of authorisation of new indication, pharmaceutical form or route of administration: not applicable		
Please note that development of the medicinal product above in the following condition(s)/indication(s):		
Treatment of chronic idiopathic arthritis (including rheumatoid arthritis, psoriatic arthritis, ankylosing spondylarthritis and juvenile idiopathic arthritis)		
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)		
$oxed{oxed}$ (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		

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

(please specify:

(please specify:

other regulatory action

other reason

Assessment of efficacy and safety data from the ContRAst programme is ongoing, however the limited efficacy demonstrated does not support a suitable benefit/risk profile for otilimab as a potential treatment for RA. As a result, GSK has decided not to progress with regulatory submissions.

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) (e.g. suspension, revocation of M.A.)

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 No		
Name and signature of the PIP contact point:	Signature on file	
Date:	25 November 2022	
Contact for inquiries from interested parties:	GlaxoSmithKline Trading Services Limited	
Telephone:	+1-438-899-8201	
Fmail:	eu.paediatric-plans@gsk.com	