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

Actives substances(s): dolute	gravir / HIV-1 maturat	ion inhibitor (GSK3640254)
Latest Decision number(s):	1) P/0523/2022	
Corresponding PIP number(s):	1) EMEA-003152-PIPO	01-21
the requirements of Article 7 or respective application) and a m there is a legal obligation to co	f the Paediatric Regular narketing authorisation mplete that PIP. The se the PIP was included	authorisation application in order to comply with tion (as a condition of the validation of the was granted based on this application, then ame applies if there has been a successful postin order to comply with the requirements of
Please confirm if any of the abo	ove applies:	
Yes □ No ⊠		
or the successful post-authoris That obligation cannot be cance must be completed, unless it is measures or granting a full pro with the Paediatric Regulation)	ation application, as apelled by a unilateral destination of a modified in agreement of a language o	sation obtained at the end of that initial procedure oplicable, you are obliged to complete that PIP. ecision, including by withdrawing the MA. Such PI with the PDCO by removing all outstanding PIP stead (upon relevant circumstances in accordance binding PIP establishes noncompliance with the European Medicines Agency has an obligation to
Please note that development condition(s)/indication(s):	of the medicinal produc	ct above in the following
Treatment of human immunode	eficiency virus (HIV-1)	infection
$oxed{\boxtimes}$ has been discontinued		
for the following reason(s): (tio	ck all that apply)	
$\hfill\Box$ (possible) lack of efficacy in	adults	
$\hfill\Box$ (possible) lack of efficacy in	children	
☐ (possible) unsatisfactory sat	fety profile in adults	
☐ (possible) unsatisfactory sat	fety profile in children	
$oxed{\boxtimes}$ commercial reasons (please	specify: Applicant dec	cision not to continue development of product)
☐ manufacturing / quality pro	blems	
$\hfill \square$ other regulatory action	(please specify:	)
other reason	(please specify:	)

Please add a brief	description	(max 2000	characters)	of the	reason(s)	for the	discontinuation:
i icasc add a bilci	ucscription	IIIIax Zuuu	Cital actors	OI LIIC	i Casoni (s	, 101 111	uiscontinuation.

Applicant has decided to discontinue development of oral GSK3640254/DTG fixed-dose combination (FDC) for commercial reasons.

Date: 11 August 2025

Contact for inquiries from interested parties: EU Paediatric Plans

Telephone: +14388998201

Email: <u>eu.paediatric-plans@gsk.com</u>

 $<sup>^{\</sup>mathrm{i}}$  This form will be published to the corresponding decision available on the website of the European Medicines Agency.