

Apixaban Accord

Procedural steps taken and scientific information after the authorisation

Application number	Scope	Opinion/ Notification ¹ issued on	Commission Decision Issued ² / amended on	Product Information affected ³	Summary
IB/0003	C.I.3.z - Change(s) in the SPC, Labelling or PL intended to implement the outcome of a procedure concerning PSUR or PASS or the outcome of the assessment done under A 45/46 - Other variation	18/05/2022		SmPC and PL	To update section 4.8 of the SmPC and section 4 of the PL to implement the wording agreed by the CHMP related to the adverse reaction vasculitis, with a frequency 'not known'.

² A Commission decision (CD) is issued for procedures that affect the terms of the marketing authorisation (e.g. summary of product characteristics, annex II, labelling, package leaflet). The CD is issued within two months of the opinion for variations falling under the scope of Article 23.1a(a) of Regulation (EU) No. 712/2012, or within one year for other procedures.

³ SmPC (Summary of Product Characteristics), Annex II, Labelling, PL (Package Leaflet).



¹ Notifications are issued for type I variations and Article 61(3) notifications (unless part of a group including a type II variation or extension application or a worksharing application). Opinions are issued for all other procedures.

IB/0001/G	This was an application for a group of variations.	01/07/2021	SmPC and PL	
	C.I.2.a - Change in the SPC, Labelling or PL of a			
	generic/hybrid/biosimilar products following			
	assessment of the same change for the reference			
	product - Implementation of change(s) for which NO			
	new additional data is required to be submitted by			
	the MAH			
	C.I.2.a - Change in the SPC, Labelling or PL of a			
	generic/hybrid/biosimilar products following			
	assessment of the same change for the reference			
	product - Implementation of change(s) for which NO			
	new additional data is required to be submitted by			
	the MAH			
	C.I.2.a - Change in the SPC, Labelling or PL of a			
	generic/hybrid/biosimilar products following			
	assessment of the same change for the reference			
	product - Implementation of change(s) for which NO			
	new additional data is required to be submitted by			
	the MAH			
	C.I.2.a - Change in the SPC, Labelling or PL of a			
	generic/hybrid/biosimilar products following			
	assessment of the same change for the reference			
	product - Implementation of change(s) for which NO			
	new additional data is required to be submitted by			
	the MAH			