

Post Authorisation Assessments

Carprieve Tablets 20 mg Vm 02000/4220

	10 Januar : 0004	Other changes to the active cubstance. Outpataintial
•	19 January 2024	Other changes to the active substance: - Substantial changes in the updated version of the ASMF or the active substance part of the dossier.
	06 June 2023	Change to comply with an update of the relevant
•	00 Julie 2023	monograph of the Ph. Eur. or national pharmacopoeia of
		a Member State.
	28 October 2022	Change in distributor details from Norbrook Laboratories
•		(GB) Ltd, 1 Saxon Way East, Oakley Hay Industrial
		Estate, Corby, Northamptonshire, NN18 9EX, United
		Kingdom to Norbrook Laboratories Limited, Carnbane
		Industrial Estate, Newry, Co. Down, BT35 6QQ, Northern
		Ireland.
•	17 March 2021	Submission of an updated Ph. Eur. certificate of
		suitability for an active substance from an already
		approved manufacturer.
•	21 December 2020	Addition of a manufacturer of the active substance or
		addition of a site of manufacture.
•	22 November 2019	Submission of a new Ph. Eur. certificate of suitability for
	45 August 2040	an active substance excipient from a new manufacturer.
•	15 August 2019	Replacement or addition of a supplier of packaging components or devices.
		Minor changes to an approved test procedure of the
		finished product.
		Increase in batch size (including batch size range) of the
		finished product.
		Decrease in batch size range of the finished product.
		Minor change in the manufacturing process of the
		finished product.
		Update of the dossier to comply with the provisions of an
		updated general monograph of the Ph. Eur for the
	04 May 2040	finished product.
•	21 May 2019	Change in the manufacturer of a starting material used in
		the manufacturing process of the active substance. Change of specification of the finished product
•	02 February 2011	
•	28 January 2009	Change of name of the product from 'Norocarp Tablets 20mg' to 'Carprieve Tablets 20mg'
•	08 August 2008	Renewal
	05 October 2006	Change in specification of the finished product
	21 September 2006	Changes to the SPC and Product Literature to bring in
•		line with new legislation
•	12 July 2006	Addition of indication regarding post-operative pain
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•	01 June 2006	Change of shelf life from 18 months (Tubs) and 12 months (Blister) to 26 months (Tubs) and 24 months (Blister)
•	12 December 2003	Additional presentation (blister pack)