

Post Authorisation Assessments

Downland Fluke and Worm Oral Suspension

Vm 02000/4169

	04 Mars 0004	
•	04 May 2024	Container closure system dimensions corrected.
		Change in dimensions on the container closure system.
		Change in dimensions on the container closure system.
•	15 September 2023	Change to comply with an update of the relevant
		monograph of the Ph. Eur. Or national pharmacopoeia of
		a member state.
•	04 May 2023	Update to Section 4.6 of the SPC and corresponding
		section of the PL.
•	10 March 2023	Editorial changes to part 2 of the dossier.
•	03 January 2023	Change of Distributor address 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, BT35 6QQ, Co
		Down, Northern Ireland.
•	21 June 2022	Change in the name and address details of an active
		substance master file (ASMF) holder.
•	13 June 2022	Change of specification(s) of a former non
		Pharmacopoeial active substance to comply with the Ph.
		Eur. or with a national pharmacopoeia of a Member
		State.
•	23 December 2021	Change in the SPC, Labelling / Package Leaflet intended
		to implement the outcome of a procedure concerning
		PSUR.
•	05 November 2020	Change in distributor details from Downland Marketing
		Limited, 15 Victoria Place, Carlisle, CA1 1EW to
		Downland Marketing Ltd, Main Mill, Warwick Mill
		Business Centre, Warwick Bridge, Carlisle, CA4 8RR.
•	24 October 2019	Tightening of specification limits of an excipient.
		Addition of a new specification parameter to the
		specification with its corresponding test method of an
		excipient.
		Deletion of a non-significant specification parameter of
		an excipient.
		Change in the specification parameters and/or limits of
		an excipient.
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		an excipient.
•	30 July 2019	Change in the QPPV of an existing pharmacovigilance
		system as described in the DDPS.
		Change of the back-up procedure of the QPPV of an
		existing pharmacovigilance system as described in the

		DDPS.
•	29 August 2018	Deletion of a supplier of packaging components or devices.
		Minor changes to an approved test procedure of the finished product.
		Minor changes to an approved test procedure of the finished product.
		Update of the test procedure to comply with the updated general Ph. Eur monograph.
		Increase in batch size (including batch size range) of the finished product.
•	28 March 2017	Deletion of a non-significant specification parameter of the finished product.
•	10 November 2014	Changes to an existing pharmacovigilance system as described in the DDPS.
•	01 September 2010	Change of withdrawal period for meat from sheep from 28 days to 5 days
•	09 June 2010	Update to SPC and Product Literature to include standard warnings for anthelmintics
•	04 September 2008	Changes to the SPC and Product Literature to bring in line with new legislation
•	07 March 2007	Change of legal category from PML to POM-VPS
•	10 April 2006	Renewal
•	12 October 2005	Addition of a manufacturing site of assembly
•	05 August 2005	Change in formulation
•	22 December 2003	Renewal
•	24 October 2003	Addition of manufacturer and assembler of the dosage form
•	13 December 2002	Addition of manufacturers of the active substance