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## **Post Authorisation Assessments**

## Prellim 0.075 mg/ml Solution for Injection for Cattle and Pigs Vm 31592/3002

•	13 February 2024	Change in name of manufacturer of the finished
•	21 April 2023	product. Change in name of manufacturer of finished product
		responsible for batch release.
		Change in MAH name and address from Laboratorios
		Syva S.A.U., Avda. Párroco Pablo Díez, 49-57, 24010
		LEÓN, ESPAÑA to Laboratorios SYVA S.A., C/
		Marqués de la Ensenada, 16, 28004 Madrid, Spain.
•	21 April 2023	Change in name of manufacturer of the finished product.
•	21 April 2023	One-off alignment of the product information with
	07.4 10000	version 9.0* of the QRD templates.
•	07 April 2022	Changes in the SPC, Labelling or Package Leaflet for
		harmonization of the SPC between original and new
	07.4	concerned Member States.
•	07 April 2022	Deletion of a pack size(s) of the finished product.
•	08 March 2022	Variation to update the Active Substance Master File.
•	13 October 2021	Increase in batch size (from 50 L batch size to batch
		range of 50 L - 157.5 L) of the finished product.
•	20 November 2020	Change in distributor details from Zoetis UK Limited,
		5th Floor, 6 St. Andrew Street London EC4A 3AE to
		Zoetis UK Limited, 1st Floor, Birchwood Building,
	17 September 2019	Springfield Drive, Leatherhead Surrey KT22 7LP. Submission of an updated ASMF for an approved
•		manufacturer of the active substance.
•	15 November 2018	Change in the number of units (e.g. tablets, ampoules,
		etc.) in a pack within the range of the currently
		approved pack sizes of the finished product.
•	10 April 2018	Repeat Use application to add 4 new member states
•	03 January 2018	Change to a test procedure for the finished product.
•	10 May 2017	Updates to the approved Active Substance Master
		File.
•	21 January 2016	Deletion of a manufacturing site.
•	06 October 2015	Addition of a specification test for the active substance.
		Addition of a new active substance manufacturer.
•	08 January 2015	Change to a limit test to comply with a Ph. Eur.
		monograph.
•	12 December 2014	Change in test procedure for the finished product.
•	23 April 2014	Renewal.
•	11 December 2013	Change of distributor.
•	28 August 2013	Change in the invented name of the product from

		'Luteosyl 0.075 mg/ml Solution for Injection' to 'Prellim 0.075 mg/ml Solution for Injection'.
•	26 August 2011	To change the manufacturer of the finished product responsible for batch release.
•	26 August 2011	To change the manufacturer of the finished product.
•	26 August 2011	To change the name of the MAH from from Laboratorios SYVA, S.A. to Laboratorios SYVA, S.A.U.