## VPA22622/022/001

## Vetivex 11 (Hartmann's) solution for infusion for cattle, horses, dogs and cats

Variation	Summary	Date
Vet - F.II.e.4 b)	VRA-R - Vet - F.II.e.4 b) - b) Sterile medicinal products - F.II.e.4 b) Quality Changes - Container closure system - Change in shape or dimensions of the container or closure (immediate packaging) - Sterile medicinal products	27/02/24
Vet - B43	VNRA - Vet - B43 - Editorial changes to part 2 of the dossier if inclusion in an upcoming procedure concerning part 2 is not possible - B43 Changes to the quality part of the dossier: Editorial changes to part 2 of the dossier if inclusion in an upcoming procedure concerning part 2 is not possible	19/09/23
Vet - F.II.e.2 z)	VRA-R - Vet - F.II.e.2 z) - z) Other changes under this code level e.g. variations outlined in section 6 and 7 of EMA/CMDv/7381/2021 - F.II.e.2 z) Quality Changes - Container closure system - Change in the specification parameters and/or limits of the immediate packaging of the finished product - Other changes under this code level, e.g. variations outlined in section 6 and 7 of EMA/CMDv/7381/2021	28/06/23
Vet - F.II.e.4 b)	VRA-R - Vet - F.II.e.4 b) - b) Sterile medicinal products - F.II.e.4 b) Quality Changes - Container closure system - Change in shape or dimensions of the container or closure (immediate packaging) - Sterile medicinal products	28/06/23
Vet - F.I.b.1 g)	VRA-R - Vet - F.I.b.1 g) - g) Change in the testing frequency of specification parameter, from routine testing to skip or periodic testing - F.I.b.1 g) Quality Changes - Active Substance - Control of active substance - Change in the specification parameters and/or limits of an active substance, starting material/intermediate/reagent used in the manufacturing process of the active substance - Change in the testing frequency of specification parameter, from routine testing to skip or periodic testing	28/06/23
Vet - F.III.1 a) 1.	VRA-R - Vet - F.III.1 a) 1 a) European Pharmacopoeial Certificate of Suitability to the relevant Ph. Eur. Monograph. 1. New certificate for a non-sterile active substance that is to be used in a sterile medicinal product, where water is used in the last steps of the synthesis and the material is not claimed to be endotoxin free - F.III.1 a) 1. Quality Changes - CEP/TSE/MONOGRAPHS - Submission of a new or updated Ph. Eur. certificate of suitability or deletion of Ph. Eur. certificate of suitability: -For an active substance -For a starting material/reagent/intermediate used in the manufacturing process of the active substance -For an excipient European Pharmacopoeial Certificate of Suitability to the relevant Ph. Eur. Monograph - New certificate for a non-sterile active substance that is to be used in a sterile	28/06/23

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	medicinal product, where water is used in the last steps of the synthesis and the material is not claimed to be endotoxin free	
Vet - F.III.1 a) 1.	VRA-R - Vet - F.III.1 a) 1 a) European Pharmacopoeial Certificate of Suitability to the relevant Ph. Eur. Monograph. 1. New certificate for a non-sterile active substance that is to be used in a sterile medicinal product, where water is used in the last steps of the synthesis and the material is not claimed to be endotoxin free - F.III.1 a) 1. Quality Changes - CEP/TSE/MONOGRAPHS - Submission of a new or updated Ph. Eur. certificate of suitability or deletion of Ph. Eur. certificate of suitability: -For an active substance -For a starting material/reagent/intermediate used in the manufacturing process of the active substance -For an excipient European Pharmacopoeial Certificate of Suitability to the relevant Ph. Eur. Monograph - New certificate for a non-sterile active substance that is to be used in a sterile medicinal product, where water is used in the last steps of the synthesis and the material is not claimed to be endotoxin free	28/06/23
Vet - F.III.1 a) 1.	VRA-R - Vet - F.III.1 a) 1 a) European Pharmacopoeial Certificate of Suitability to the relevant Ph. Eur. Monograph. 1. New certificate for a non-sterile active substance that is to be used in a sterile medicinal product, where water is used in the last steps of the synthesis and the material is not claimed to be endotoxin free - F.III.1 a) 1. Quality Changes - CEP/TSE/MONOGRAPHS - Submission of a new or updated Ph. Eur. certificate of suitability or deletion of Ph. Eur. certificate of suitability: -For an active substance -For a starting material/reagent/intermediate used in the manufacturing process of the active substance -For an excipient European Pharmacopoeial Certificate of Suitability to the relevant Ph. Eur. Monograph - New certificate for a non-sterile active substance that is to be used in a sterile medicinal product, where water is used in the last steps of the synthesis and the material is not claimed to be endotoxin free	28/06/23
Vet - F.II.b.2 b) z.	VRA-R - Vet - F.II.b.2 b) z b) Replacement or addition of a manufacturer responsible for importation and/or batch release z. Other changes under this code level, e.g. variations outlined in section 6 and 7 of EMA/CMDv/7381/2021 - F.II.b.2 b) z. Quality Changes - Finished Product -Manufacture - Change to importer, batch release arrangements and quality control testing of the finished product - Replacement or addition of a manufacturer responsible for importation and/or batch release - Other changes under this code level, e.g. variations outlined in section 6 and 7 of EMA/CMDv/7381/2021	28/06/23
Vet - F.II.b.1 d)	VRA-R - Vet - F.II.b.1 d) - d) Site where any manufacturing operation(s) take place, except batch release, batch control, and secondary packaging, for sterile veterianry medicinal products (including those that are aseptically manufactured) excluding biological/ immunological veterinary medicinal products - F.II.b.1 d) Quality Changes - Finished Product	28/06/23

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	-Manufacture - Replacement or addition of a manufacturing	
	site for part or all of the manufacturing process of the finished	
	product - Site where any manufacturing operation(s) take	
	place, except batch release, batch control, and secondary	
	packaging, for sterile veterianry medicinal products (including	
	those that are aseptically manufactured) excluding biological/	
	immunological veterinary medicinal products	
	VNRA - Vet - B47 b) - b) Change to comply with an update of	
	the relevant monograph of the Ph. Eur. or national	
	pharmacopoeia of a Member State - B47 b) Changes to the	
Vet - B47 b)	quality part of the dossier: Change to comply with Ph. Eur. or	27/06/23
	with a national pharmacopoeia of a Member State: — change	
	to comply with an update of the relevant monograph of the Ph.	
	Eur. or national pharmacopoeia of a Member State	
	VNRA - Vet - B21 - Replacement or addition of a secondary	
	packaging site of a finished product - B21 Changes to the	
Vet - B21	quality part of the dossier: Replacement or addition of a	27/06/23
	secondary packaging site of a finished product	
	VNRA - Vet - B47 b) - b) Change to comply with an update of	
	the relevant monograph of the Ph. Eur. or national	
	pharmacopoeia of a Member State - B47 b) Changes to the	
$\mathbf{V}_{at} = \mathbf{D} \mathbf{A} 7 \mathbf{b}$		27/06/22
Vet - B47 b)	quality part of the dossier: Change to comply with Ph. Eur. or	27/06/23
	with a national pharmacopoeia of a Member State: — change	
	to comply with an update of the relevant monograph of the Ph.	
	Eur. or national pharmacopoeia of a Member State	
	VNRA - Vet - B43 - Editorial changes to part 2 of the dossier	
	if inclusion in an upcoming procedure concerning part 2 is not	/ /
Vet - B43	possible - B43 Changes to the quality part of the dossier:	27/06/23
	Editorial changes to part 2 of the dossier if inclusion in an	
	upcoming procedure concerning part 2 is not possible	
	VNRA - Vet - B22 - Change to importer, batch control	
	arrangements and quality testing (replacement or addition of a	
Vot D22	site) for a finished product - B22 Changes to the quality part	27/06/22
Vet - B22	of the dossier: Change to importer, batch control arrangements	27/06/23
	and quality testing (replacement or addition of a site) for a	
	finished product	
	VNRA - Vet - B38 - Change in pack size (number of units e.g.	
	tablets, ampoules, etc. in a pack) within the range of the	
	currently approved pack size - B38 Changes to the quality part	
	of the dossier: Change in pack size (number of units e.g.	
	tablets, ampoules, etc. in a pack) within the range of the	
Vet - B38	currently approved pack size. In cases where a given pack size	26/06/23
	has received an individual marketing authorisation which is	20/00/23
	separate to the marketing authorisation for other pack sizes of	
	the same product, the change of the former will not be a	
	variation according to Article 61, but a variation according to	
	Article 62 of Regulation (EU) 2019/6	
	VNRA - Vet - B38 - Change in pack size (number of units e.g.	
Vet - B38	tablets, ampoules, etc. in a pack) within the range of the	26/06/23
	currently approved pack size - B38 Changes to the quality part	

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	of the dossier: Change in pack size (number of units e.g.	
	tablets, ampoules, etc. in a pack) within the range of the	
	currently approved pack size. In cases where a given pack size	
	has received an individual marketing authorisation which is	
	separate to the marketing authorisation for other pack sizes of	
	the same product, the change of the former will not be a	
	variation according to Article 61, but a variation according to	
	Article 62 of Regulation (EU) 2019/6	
	VNRA - Vet - B47 b) - b) Change to comply with an update of	
	the relevant monograph of the Ph. Eur. or national	
	pharmacopoeia of a Member State - B47 b) Changes to the	
Vet - B47 b)	quality part of the dossier: Change to comply with Ph. Eur. or	11/01/23
	with a national pharmacopoeia of a Member State: — change	
	to comply with an update of the relevant monograph of the Ph.	
	Eur. or national pharmacopoeia of a Member State	
	VNRA - Vet - B47 b) - b) Change to comply with an update of	
	the relevant monograph of the Ph. Eur. or national	
	pharmacopoeia of a Member State - B47 b) Changes to the	
Vet - B47 b)	quality part of the dossier: Change to comply with Ph. Eur. or	11/01/23
	with a national pharmacopoeia of a Member State: — change	
	to comply with an update of the relevant monograph of the Ph.	
	Eur. or national pharmacopoeia of a Member State	
	VNRA - Vet - B45 - Submission of a new Ph. Eur. CEP from a	
	new manufacturer (replacement or addition) for a non-sterile	
	active substance, starting material, reagent or intermediate,	
	excipient - B45 Changes to the quality part of the dossier:	
Vet - B45	Submission of a new Ph. Eur. CEP from a new manufacturer	11/01/23
	(replacement or addition) for a non-sterile: — active	
	substance; — starting material, reagent or intermediate used in	
	the manufacturing process of the active substance; —	
	excipient	
	VNRA - Vet - B47 b) - b) Change to comply with an update of	
	the relevant monograph of the Ph. Eur. or national	
Vet - B47 b)	pharmacopoeia of a Member State - B47 b) Changes to the	
	quality part of the dossier: Change to comply with Ph. Eur. or	21/12/22
	with a national pharmacopoeia of a Member State: — change	
	to comply with an update of the relevant monograph of the Ph.	
	Eur. or national pharmacopoeia of a Member State	