VPA22020/013/001

Atopica 100 mg/ml oral solution for cats and dogs

Variation	Summary	Date
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	05/01/24
	changes to part 2 of the dossier if inclusion in an upcoming procedure concerning part 2 is not possible	
Vet - B11 b)	VNRA - Vet - B11 b) - b) Tightening of specification limits of an active substance, starting material, intermediate or reagent used in the manufacturing process of the active substance - B11 b) Changes to the quality part of the dossier: Change in the specification parameters or limits of an active substance, starting material, intermediate or reagent used in the manufacturing process of the active substance or of the immediate packaging of the active substance — tightening of specification limits of an active substance, starting material, intermediate or reagent used in the manufacturing process of the active substance	30/11/23
Vet - B11 b)	VNRA - Vet - B11 b) - b) Tightening of specification limits of an active substance, starting material, intermediate or reagent used in the manufacturing process of the active substance - B11 b) Changes to the quality part of the dossier: Change in the specification parameters or limits of an active substance, starting material, intermediate or reagent used in the manufacturing process of the active substance or of the immediate packaging of the active substance — tightening of specification limits of an active substance, starting material, intermediate or reagent used in the manufacturing process of the active substance	30/11/23
Vet - B11 b)	VNRA - Vet - B11 b) - b) Tightening of specification limits of an active substance, starting material, intermediate or reagent used in the manufacturing process of the active substance - B11 b) Changes to the quality part of the dossier: Change in the specification parameters or limits of an active substance, starting material, intermediate or reagent used in the manufacturing process of the active substance or of the immediate packaging of the active substance — tightening of specification limits of an active substance, starting material, intermediate or reagent used in the manufacturing process of the active substance	30/11/23
Vet - B11 b)	VNRA - Vet - B11 b) - b) Tightening of specification limits of an active substance, starting material, intermediate or reagent used in the manufacturing process of the active substance - B11 b) Changes to the quality part of the dossier: Change in the specification parameters or limits of an active substance, starting material, intermediate or reagent used in the manufacturing process of the active substance or of the immediate packaging of the active substance — tightening of specification limits of an active substance, starting material, intermediate or reagent used in	30/11/23

	the manufacturing process of the active substance	
	VNRA - Vet - B12 a) - a) Minor changes to an approved test	
Vet - B12 a)	procedure (active, finished product, packaging, measuiring device)	
	- B12 a) Changes to the quality part of the dossier: Minor changes	
	— to an approved test procedure — for active substance; — for	18/10/23
	the finished product; —for the immediate packaging of the active	10/10/23
	substance or the finished product; — of a measuring or	
	administration device	
	VNRA - Vet - B12 a) - a) Minor changes to an approved test	
Vet - B12 a)	procedure (active, finished product, packaging, measuiring device)	
	- B12 a) Changes to the quality part of the dossier: Minor changes	
	— to an approved test procedure — for active substance; — for	18/10/23
	the finished product; —for the immediate packaging of the active	10/10/23
	substance or the finished product; — of a measuring or	
	administration device	
	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	18/10/23
	(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 pharmacopoeia of	
Vet - B47 b)	a Member State - B47 b) Changes to the quality part of the	
	dossier: Change to comply with Ph. Eur. or with a national	18/10/23
	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 - A1 b) - b) Change in the name or address or contact	
	details of a manufacturer or supplier of the active substance,	
Vet - A1 b)	starting material, reagent or intermediate used in the manufacture	İ
	of the active substance or a quality control testing site (where	
	ecified in the dossier) where no European Pharmacopoeia (Ph.	
	Eur.) Certificate of Suitability (CEP) is part of the approved	22/09/23
	dossier A1 b) Administratvie changes: Change in the name or	
	address or contact details of a manufacturer or supplier of the	
	active substance, starting material, reagent or intermediate used in	
	the manufacture of the active substance or a quality control	
	testing site (where ecified in the dossier) where no European	
	Pharmacopoeia (Ph. Eur.) Certificate of Suitability (CEP) is part	
	of the approved dossier.	
Vet - C4	VNRA - Vet - C4 - Change(s) in the SPC, labelling or package	
	leaflet intended to implement the outcome of a procedure or	
	recommendation from the competent authority or the Agency	
	concerning risk management measures in pharmacovigilance	05/07/23
	related to veterinary medicinal products - C4 Changes to the	
	safety, efficacy and pharmacovigilance part of the dossier:	
	Change(s) in the SPC, labelling or package leaflet intended to	

	implement the outcome of a mage-1 1-t' C	
	implement the outcome of a procedure or recommendation from	
	the competent authority or the Agency concerning risk	
	management measures in pharmacovigilance related to veterinary	
	medicinal products	
Vet - G.I.18	VRA-S - Vet - G.I.18 - One-off alignment of the product	04/05/23
	information with version 9.0 (or the latest version of the QRD	
	templates that are in effect at the time that this one-off variation is	
	submitted) of the QRD templates i.e. major update of the QRD	
	templates in accordance with Regulation (EU) 2019/6, for	
	veterinary medicinal products placed on the market in accordance	
	with Directive 2001/82/EC or Regulation (EC) No 726/2004 -	
	G.I.18 Safety, Efficacy, Pharmacovigilance changes - One-off	
	alignment of the product information with version 9.0 (or the	
	latest version of the QRD templates that are in effect at the time	
	that this one-off variation is submitted) of the QRD templates i.e.	
	major update of the QRD templates in accordance with	
	Regulation (EU) 2019/6, for veterinary medicinal products placed	
	on the market in accordance with Directive 2001/82/EC or	
	Regulation (EC) No 726/2004	
	VRA-R - Vet - F.I.d.1 c) - c) Extension or introduction of a re-test	09/01/23
	period/storage period supported by real time data - F.I.d.1 c)	
Vet - F.I.d.1 c)	Quality Changes - Active Substance - Stability - Change in the	
	re-test period/storage period of the active substance where no Ph.	
	Eur. Certificate of Suitability covering the retest period is part of	
	the approved dossier - Extension or introduction of a re-test	
	period/storage period supported by real time data	
	IA - A.4 - A.4 Change in the name and/or address of: a	
A.4	manufacturer (including where relevant quality control testing	
	sites); or an ASMF holder; or a supplier of the active substance,	
	starting material, reagent or intermediate used in the manufacture	11/02/22
	of the active substance (where specified in the technical dossier)	
	where no Ph. Eur. Certificate of Suitability is part of the approved	
	dossier; or a manufacturer of a novel excipient (where specified	
	in the technical dossier) - A.4 - ADMINISTRATIVE CHANGES	
	- Change in the name and/or address of: a manufacturer	
	(including where relevant quality control testing sites); or an	
	ASMF holder; or a supplier of the active substance, starting	
	material, reagent or intermediate used in the manufacture of the	
	active substance (where specified in the technical dossier) where	
	no Ph. Eur. Certificate of Suitability is part of the approved	
	dossier; or a manufacturer of a novel excipient (where specified	
	in the technical dossier)	