



Pharmacovigilance vétérinaire Evaluation du lien de causalité entre un évènement indésirable et un médicament : le système ABON

L'évaluation du lien de causalité entre un évènement indésirable déclaré et le(s) médicament(s) concerné(s) n'est plus obligatoire depuis l'entrée en vigueur du règlement 2019/6 CE.

En France, l'ANMV et le CPVL ont fait le choix de poursuivre cette évaluation pour chaque déclaration de pharmacovigilance. En effet, la conclusion de cette évaluation est transmise au déclarant qui envoie directement sa déclaration de pharmacovigilance à l'ANMV ou au CPVL.

De plus, l'évaluation du lien de causalité de chaque déclaration enregistrée dans la base nationale de pharmacovigilance permet également à l'ANMV de prioriser les signaux qui peuvent être détectés dans le cadre de la détection de signal qu'elle a mis en place.

Ce choix de maintenir cette évaluation a également été retenu par d'autres autorités compétentes nationales ainsi que par la plupart des titulaires d'AMM.

La méthodologie d'évaluation utilisée par l'ANMV et le CPVL reste celle qui était utilisée avant le règlement 2019/6 et qui est préconisée par le système ABON.

Selon ce système, il existe cinq catégories de causalité :

- Catégorie A : Probable.
- Catégorie B : Possible.
- Catégorie O : Non classable/non évaluable (effets/évènements pour lesquels les informations disponibles sont insuffisantes pour tirer des conclusions).
- Catégorie O1 : Non concluant (effets/évènements pour lesquels d'autres facteurs ont empêché de tirer des conclusions, mais une association avec le produit ne peut pas être exclue).
- Catégorie N : Improbable.

Pour l'évaluation du lien de causalité, il convient de prendre en compte les facteurs suivants :

- Lien d'association en fonction du temps incluant le dechallenge suite à l'arrêt du traitement et le rechallenge suite à une administration répétée (dans les antécédents cliniques) ou en fonction des sites anatomiques.
- Explication pharmacologique, taux sanguins, connaissance antérieure du médicament.
- Présence de phénomènes cliniques ou pathologiques caractéristiques.
- Exclusion d'autres causes.
- Exhaustivité et fiabilité des données de la déclaration.
- Mesure quantitative du degré de contribution d'un médicament vétérinaire au développement d'un effet/évènement indésirable (relation dose-effet).





Pour l'inclusion dans la catégorie « A » (probable), les critères minimaux suivants doivent être remplis :

La chronologie, entre l'administration du médicament vétérinaire et, l'apparition et la durée de l'effet/évènement indésirable signalé, est compatible.

La description des phénomènes cliniques est compatible, ou tout au moins plausible, compte tenu de la pharmacologie et toxicologie connues du médicament.

Il ne doit pas y avoir d'autre explication tout aussi plausible du cas (si d'autres explications sont suggérées, sont-elles valables? Quel est leur degré de certitude?). En particulier, l'usage concomitant d'autres médicaments vétérinaires (et d'éventuelles interactions) ou une maladie intercurrente doivent être pris en considération dans l'évaluation.

Si l'un des critères ci-dessus n'est pas rempli (en raison de données contradictoires ou par manque d'information), alors la déclaration ne peut être classée que dans la catégorie « B » (possible), « N » (improbable), « O1 » (non concluant) ou « O » (non classable/non évaluable).

Inclusion dans la catégorie « B » (possible): il est recommandé de choisir cette catégorie lorsque la causalité d'un médicament vétérinaire est l'une des causes (parmi d'autres causes) possibles et plausibles de l'effet/évènement indésirable décrit mais que les données ne permettent pas de satisfaire le critère d'inclusion dans la catégorie « A ».

Inclusion dans la catégorie « O » (non classable / non évaluable): tous les cas pour lesquels on ne dispose pas de données fiables suffisantes, voir d'aucune donnée fiable, concernant un effet/évènement indésirable pour pouvoir évaluer le lien de causalité.

Inclusion dans la catégorie « O1 » (non concluant): tous les cas pour lesquels un lien avec le médicament vétérinaire ne peut pas être exclu mais pour lesquels d'autres facteurs empêchent de tirer des conclusions.

Inclusion dans la catégorie « N » (improbable): les cas pour lesquels il existe suffisamment de données pour établir hors de tout doute raisonnable qu'un autre facteur est à l'origine de l'effet/évènement indésirable sans aucun rapport avec le médicament vétérinaire.

D'autres orientations sur la façon d'évaluer le lien de causalité sont disponibles dans l'ancienne ligne directrice du CVMP sur l'harmonisation de l'approche de l'évaluation du lien de causalité pour les effets indésirables des médicaments vétérinaires (CVMP Guideline on Harmonising the Approach to Causality Assessment for Adverse Reactions to Veterinary Medicinal Products) et un extrait est rappelé en annexe.





Annexes

Recommendation on harmonising the approach to causality assessment for adverse events to veterinary medicinal products (extract)

4. Questionnaire

4.1. Associative connection

- a. in time (including de-challenge and re-challenge)
- b. with anatomical site.

4.1.1.

Is the observed event associated with the administration of the VMP? Is the chronology in good accordance with treatment? Is there a reasonable association in time between the administration of the product and the onset and duration of the adverse event?

• Is there a reasonable association in time between the administration of the product and the onset of the adverse event?

yes	no	not known
reasonable association	no reasonable association	unknown
A, B	N	O1 or O

4.1.2.

Has there been any	no	not known
improvement	no improvement	no de-challenge done
А, В	O, N	A, B, O1, O, N

4.1.3.

Did the adverse event reappear after re-challenge (same or related animal)? Is a similar event known in that patient from previous exposure?

• What happened after re-challenge - recurrence, no recurrence or no re-challenge done?

yes	no	not known
recurrence	no recurrence	no re-challenge done
А. В	N	A. B. O1. O. N

4.1.4.

Could the location/distribution of signs be caused by the treatment?

yes	no	not applicable
associative anatomical	no anatomical connection	
connection		
А, В	N	A, B, O1, O, N



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Main question for section 4.1

• Is there a reasonable association in time and/or anatomical site?

yes	no	not known
reasonable association	no reasonable association	unknown
A, B	N	O1 or O

4.2. Pharmacological and/or immunological explanation

- known pharmacology, toxicology of the product (active substance and/or excipients)
- VMP concentrations in blood
- dose-effect relationship (degree of contribution of a product to the development of a reaction).

4.2.1.

Does the reported event fit into the toxicological profile or allergic potential of the product? Does the pharmacological/toxicological knowledge of the product fit the signs? Is the adverse event, the description of the clinical phenomena, consistent with or at least plausible, given the known pharmacology and toxicology of the product?

Do similar compounds cause events of this type?

• Does the reported event fit into the pharmacological/toxicological profile or allergic potential of the product?

yes	no	
A, B	01, O, N	

4.2.2.

Has the product been overdosed? Did the product concentration in blood exceed the therapeutic concentration? Are concentrations in plasma known? What dose was used - overdose, correct dose, low dose, unknown dose? Did the adverse event show a dose-effect relationship?

Did the adverse event show a dose-effect relationship (e.g. overdose)?

yes	no	not known
А, В	A, B, O1, O, N	A, B, O1, O, N

Main question for section 4.2

• Is there a reasonable association with the known pharmacological/toxicological profile, the allergic potential of the product and/or a dose-effect relation?

yes	no	
A, B	01, O, N	





4.3. Presence of characteristic product or treatment related clinical or pathological phenomena

Are characteristic clinical or pathological phenomena present, which are related to the product or treatment?

Are there any measurable criteria to confirm the adverse event objectively, are confirming factors known (post mortem results, laboratory results)?

• Are additional data (laboratory tests, pathological findings) confirming clinical plausibility?

yes	no	not applicable/not available
A, B	N	A, B, O1, O, N

4.4. Previous knowledge of similar reports

- a. from literature
- b. from adverse events reported before

Are there any reports of this event known from literature? Is the event known and expected (described in SPC)? Have there been previous reports with these kinds of signs? Was this type of event reported before in an adverse event? Is the adverse event (generally) known to be potentially related to the product or treatment mentioned? ('adverse event' in this respect is the single pathological sign or the [majority of the signs in the] complex. 'Known' means published in literature or reported before and classified as A (probable) or B (possible)).

• What about consistency of the reported event - is it already described in literature or SPC, has it been reported before?

yes	yes	no	no
described in literature	observed before,	never observed before,	never observed before,
or	but not fitting	but fitting pharm./tox.	not fitting pharm./tox.
SPC, described in case	pharm./tox. profile	profile	profile
A, B	B, O1, O, N	B, O1, O, N	Ó1, O, N

4.5. Exclusion of other causes

Are there possible other causes for the adverse event? Is there another (also) likely cause? Is there another obviously more likely cause? Is this adverse event, to my best knowledge, unrelated to treatment? Use of combination of products/other products used?

Is the present disease contributing to signs? Is the health status of the animal contributing to signs? Are predisposing factors known? Are there other confirmed causes known (post mortem results, laboratory results, re-/de-challenge, other products used with pharmacological-toxicological potential to cause this event)?

Is there any other explanation (confirmed, possible, no other explanation)?

yes	yes	no
confirmed	possible	none
N	B, O1, O	Α





4.6. Completeness and reliability of the data in the case reports

 Is the reported information insufficient? Is there reason to doubt the reporting source/information?

yes	no	
01, 0	A, B, N	

5. Causality assessment by judging the answers to the questionnaire - minimum criteria

5.1. For inclusion in category A (probable)

Associative connection in time (4.1 = yes) and

Adverse event fits the pharmacological/toxicological profile of the product (4.2 = yes) and

No other equally plausible explanation (4.5 = no) and

No indication of insufficient/unreliable information (4.6 = no).

5.2. For inclusion in category B (possible)

Associative connection in time (4.1 = yes) and

Adverse event fits the pharmacological/toxicological profile of the product (4.2 = yes) and

Other equally plausible explanation possible (4.5 = yes) and

No indication of insufficient/unreliable information (4.6 = no).

or

There have been reports of the adverse event before (4.4 = yes) and No indication of insufficient/unreliable information (4.6 = no) and

Associative connection in time (4.1 = yes) or adverse event fits the pharmacological/toxicological profile of the product (4.2 = yes).

5.3. For inclusion in category O1 (inconclusive)

Category O1 is for events where at least one of the answers from the questionnaire point to a causal relationship to the product or the treatment (A or B) but overall information is not sufficient to draw a conclusion. As some of these O1 classified events will recur and due to sufficient information in subsequent reports turn out to belong to B or even A category, they present an interesting issue for surveillance. For pharmacovigilance surveillance purposes O1 classified events can be seen as kind of interesting "precursors".





Associative connection in time (4.1 = yes) and/or

Adverse event fits into the pharmacological/toxicological profile of the product (4.2 = yes) and/or

No other equally plausible explanation (4.5 = no) and

Inconclusive, unreliable or insufficient information (4.6 = yes).

5.4. For inclusion in category O (unclassifiable/unassessable)

Inconclusive, unreliable or insufficient information (4.6 = yes) which cannot be used to answer questions 4.1 to 4.5.

5.5. For inclusion in category N (unlikely)

Sufficient information exists to confirm that the product or treatment did not cause the adverse event (4.5 = yes) and

No indication of insufficient/unreliable information (4.6 = no).





Annex 1:

Examples of adverse events for the application of the causality assessment guidance

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Adverse event number	A 1	A 2	В 1	B 2	В 3	В 4
Therapeutic group	Antiemetic	Antibiotic	Antibiotic	Anaesthetic	Antibiotic	Sex hormones/ modulators of the genital system
Active ingredients Other products	Dopamine D2- antagonist	Sulfonamides	Tetracycline group Oxytocic agent	r- cotic agent α - adrenoceptor agonist, α 2-	Aminoglycosid	Antiprogestogenic agent
Species, Breed Sex, Age, Weight	Dog West Highland White Terrier 6 years	Dog	Horse	Dog, Cat	Dog, Labrador Retriever female 9 months 27 kg	Dog Shi Tzu 9 years
Nos. treated	1	1	1	2 2	1	1
Nos. died	Ö	Ö	1	0	Ö	0





ABON A A B B B B B		For gastritis, a west highland white terrier was injected with the dopamine antagonist. Rapidly, it displayed an abnormal behaviour, aggression, abnormal gait, tremors and excitation. Within a few hours, it recovered spontaneously	Kerato- conjunctivitis sicca after three weeks of treatment	Minutes after i.v.injection staggering, collapse, death	5 minutes after injection respiratory depression, abnormal breathing	24 hrs after the treatment (for eczema in one ear) the dog showed diarrhoea. At this time the dog began oestrus. The treatment was suspended and the diarrhoea disappeared in two days.	For a suspicion of pregnancy, a female Shih Tzu was injected subcutaneously with 2 ml of an antiprogestogenic agent. During the injection, the dog moved and the veterinarian suspected intravenous injection (haematoma on the injection point). A few minutes later, the dog staggered, displayed recumbency, bradycardia, dyspnoea and shock. With a symptomatic treatment (diuretic, 2 different glucocorticoids), the signs decreased within one hour. The dog displayed then anorexia, weight loss, diarrhoea and depression during 4 days.
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Adverse event number	A 1	A 2	B 1	B 2	В 3	B 4
Reasons for coding the example	A: Chronology and troubles are compatible a suspected adverse effect of the antagonist. Numerous similar reports are already registered, no equally explanation.	reaction, known suspected adverse effect, time of onset and dose fit in, no equally plausible explanation.	B: Onset of signs almost immediately following i/v administration, observed event consistent with previous reports; a second (oxytocic agent, i.v.) had administered to the animal at approx. the same time. Horse not a target species. Product past expiry - other equally plausible explanation.	but it is distinguish the product responsible the adverse event- co-	the antibiotic affected the gut flora and caused the diarrhoea. described in literature, but	





Adverse event number	01	O 2	N 1	N 2
Therapeutic group	Vaccine	NSAID	Antiparasitic (spot on)	Agents acting on the autonomic nervous system
Active ingredients Other products	Vaccine	Arylpropionic acid derivative	Avermectin	Indirect acting sympathomimetic
Species, Breed, Sex, Age, Weight	Cat	Dog 3 years	Dog Basset 8.5 years	Dog
Nos. treated	1	1	1	1
Nos. reacted Nos. died	1 0	1 0	1 0	1
Description of the event	A healthy cat becomes ataxic the day after a vaccination. No fever or other signs, and no signs of other diseases. The signs disappear after a few days.		Call before examination. The dog was treated with one pipette of the avermectin- containing product (20-40 kg). A few hours later, the dog displayed hind limb paresis. Note: the owner carried the animal to get out of the car. In fact, after examination, the animal displayed a protrusion of a vertebral disk.	Hours later halitosis, vomiting, diarrhoea, death.
ABON	O1	0	N	N





	an overall B classification. Not	totally unreliable. O	permits to exclude	N: Animal was ill prior to treatment; end stage renal disease diagnosed at post mortem
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Annex 2: Assessment of adverse events recorded as "off-label use" reports

Background

The scope of veterinary pharmacovigilance as defined in Article 73 of Directive 2001/82/EC does not only cover adverse events in animals to VMPs used under authorised conditions of use, but also any available information related to reports after off-label use of VMPs.

Reports of adverse events may be obtained on VMPs used outside the terms of the marketing authorisation e.g. use in non-authorised species, use in non-authorised species for indications which are not authorised, use at doses or via application routes differing from those set out in the relevant SPC. Such reports can provide useful information on the safety of the VMP and should be recorded and reported to the competent authorities under the responsibility of the qualified person responsible for pharmacovigilance. Periodic safety update reports should include all (serious and non-serious) reports of off-label use of the VMP.

Definition

Off-label use: the use of a VMP that is not in accordance with the SPC, including the misuse and serious abuse of the product [as defined in Article1 (16) of Directive 2001/82/EC].

Other terms often used in this context should not be used to avoid misunderstanding, e.g.:

- extra-label use
- extra-label drug use (ELDU)

Criteria to be considered when classifying adverse events as "off-label use"

In general, the use of a VMP has to be in accordance with the SPC. However, situations occur where medicinal products are used – on purpose or unintended - in a way which is not covered by the SPC. Experience from Member States shows that the frequency of under-reporting for off-label use is much higher and follow-up is more difficult to perform than for 'regular' events occurring after recommended use. Often veterinary surgeons are hesitant to report an adverse event due to the off-label-use or to give further details, which would be necessary for the comprehensive assessment of the event. One reason for this could be the fear of legal and/or financial consequences.

Off-label use reports can provide useful information on the safety of the given VMP, e.g. it can reveal risks of incorrect administration and should be recorded under the responsibility of the person responsible for pharmacovigilance and reported to the competent authorities in the same way as for all other adverse events. The existing reporting procedures should be used.

Reports of adverse events concerning off off-label use may be obtained:

- on products used outside the terms of the marketing authorisation e.g. use of a product in nonauthorised species, use at doses differing from those set out in the SPC and product information.





There are various possibilities where the use of a VMP is not in accordance with the SPC:

- Target species not authorised (special case: 'cascade', see below)
- Category or age of animal not authorised

Some products are only authorised for specific animal sub-categories e.g. a vaccine may be only recommended for active immunisation of sows and gilts or treatment is only authorised for a specific age-category.

Use during pregnancy, lactation or lay

Very often the treatment or vaccination of pregnant and lactating animals is not recommended as this has not been investigated.

- Breed not authorised
- Incorrect route of administration
- Incorrect injection site

For several products the injection site is recommended e.g. birds should be given vaccines subcutaneously into the lower part of the neck.

- o Wrong dosage or treatment scheme
- o Wrong reconstitution of the medicine

This may happen with products such as live vaccines which are reconstituted with a different diluent or another vaccine.

o Use of a VMP with an expired date. - when products are used concurrently

All medication used or administered over at least a one one-week period preceding the adverse event should be provided when available. However, a large number of VMPs, mostly vaccines, state in the SPC that no information is available on safety and efficacy when used with other products (vaccines). A decision to use the product before or after any other product therefore "needs to be made on a case to case basis". This reflects the need for collecting more information on concurrent use. It is therefore recommended to equally provide details of all medication used over at least a one week period preceding the adverse event.

At the same time it should be clear that if another product has been used concurrently, any adverse event report for a product used in line with the SPC (and according to the SPC recommendation on concurrent use had to be made on a case to case basis) will not be classified as off-label.

- on products used outside the terms of the marketing authorisation but in conformity with the provisions of Article 10 or 11 of Directive 2001/82/EC - use of unauthorised VMPs





In Articles 10 and 11 of Directive 2001/82/EC an exception is mentioned known as the prescribing "cascade". The "cascade" allows the veterinary surgeon to use products following a series of decisions providing that no authorised product is available for the treatment of that specific patient. The veterinary surgeon may prescribe:

- i. A product authorised in their Member State for that condition in another species or a product authorised for another condition but in the same species
- ii. If no such product exists, an appropriate authorised human medicine or a VMP authorised in another EU Member State
- iii. If no such product exists, a product prepared extemporaneously by an authorised person in accordance with a prescription.
- use of illegal medicines (misuse, abuse).

Causality assessment for adverse events after off-label use

To ensure consistency in using the ABON-system by pharmacovigilance personnel in competent authorities as well as in industry, common approaches to analyse, assess and code reported off-label use events should be established.

The overall assessment of off-label reports is essentially the same as for 'regular' reports (following recommended use of the VMP) and follows the rules laid down in Volume 9B resulting in a causality assessment according to the ABON scheme: A (probable), B (possible), O1 (inconclusive) or O (unclassifiable/unassessable) or N (unlikely). The guidance for causality assessment in Volume 9B does not mention the use of a product according or against the instructions for use as being relevant for the causality assessment.

However, the experience so far has revealed a tendency to classify such events as N or O causality. The fact that the product per definition is not used as recommended may suggest that these events are classified differently.

However, it has to be remembered that causality assessment takes into consideration both product and treatment: it addresses the issue of whether and how the reported treatment with the product and the reported adverse events are causally related - irrespective whether the product is used according to the recommendations for use or off-label - whereas regulatory actions will generally be triggered by at least potentially product related causality. Nevertheless, any at least possibly causally related serious off-label events where a potential risk using the product incorrectly has been identified may necessitate changes in SPC (e.g. warnings or explanation of correct use).





Annex 3:

Assessment of adverse events recorded as lack of expected efficacy (LEE) concerning pharmaceuticals

Background

Directive 2001/82/EC cites the failure to demonstrate efficacy as a reason for refusal or revocation of a marketing authorisation. It is an important aspect of the consideration of the benefit-risk balance of a product. It is felt necessary to provide a basis for a common understanding and uniformity in assessing adverse events recorded as LEE.

For the time being guidance on LEE is provided for pharmaceutical only, vaccines are excluded. Guidance on LEE concerning vaccines will be provided in due time.

Criteria to be considered when classifying adverse events as LEE

According to the definition of Volume 9B, "LEE may be defined as the apparent inability of an

authorised product to have the recognised expected efficacy in an animal, according to the claims of the SPC and following use of the product in accordance with the SPC."

It was concluded that LEE should only be considered as such when the VMP was administered according to the claims of the SPC and following use of the product in accordance with the SPC.

Pharmaceutical overdose events are usually exceptions to the requirement that qualifying an event as LEE the VMP needs to be administered according to the claims of the SPC and following use of the product in accordance with the SPC. The information related to the therapeutic indications, the route of administration, the dosage and the target species (age and all other animal characteristics data) should be checked and analysed from a critical point of view before assessing such an event which is identified as LEE by the reporters. The laboratory investigations/postmortem examination to confirm the involvement of the product or to establish a differential diagnosis are very important to thoroughly assess these events.

Events should be recorded as LEE after having been administered at a dose higher than that recommended. For instance, if a VMP administered at twice the recommended dose is not efficacious, it is reasonable to assume that it should be non efficacious when administered at the recommended dose. In certain circumstances, products used at higher doses than those recommended can give rise to cases of LEE, e.g. anthelmintic resistance on a farm.

Factors to take into account for the causality assessment of LEE reports

To ensure consistency in using the ABON-system by pharmacovigilance personnel in competent authorities as well as in industry, common approaches to analysing, assessing and coding of reported adverse events have been adopted.







Eight main factors should be taken into account: the conditions of administration, clinical or pathological signs, clinical explanation, environmental situation, onset of clinical signs, other causes, hygiene conditions, quality defect, reliability of data, in particularly the reliability of diagnosis i.e. diagnosis made by a veterinary surgeon or animal owner versus clinical diagnosis confirmed by laboratory and post mortem investigations) and published data. For LEE reports it is essential to substantiate clinical observations by laboratory data (post-mortem reports, microbiological and/or parasitological investigations). A determined effort should be made to gain additional laboratory data supporting clinical observations when the LEE report is received without this information.

Causality assessment of LEE reports

The following approach compiles guiding questions for each aspect, which are meant as examples and not intended to be exhaustive. They facilitate finding the answer to the main questions, which are listed at the end of each section (c.f. Table). According to the question and the information available, a choice of 2 or 3 answers is given: yes, no or unknown, some answers point towards N coding. The overall interpretation of the answers point towards A (probable), B (possible), O1 or O (inconclusive or unclassifiable/unassessable) or N (unlikely). In the future, an algorithm could be a useful tool to achieve consistency in causality assessment.







1. Was th	ne VMP use	ed in accordance with recommendation of th	ne marketing authorisation?		
	1.1.	1.1. Were the therapeutic indications respected? [A clear NO points towards N (unlikely)]			
		Were the characteristics of the animals to which the VMP has been administered in			
	1.2.	compliance with the SPC recommendations (species, age etc.)? [A clear NO points towards N (unlikely)]			
		Was the dose administered correct (in comp	oliance with the SPC recommendations)?		
	1.3.				
		[A clear NO points towards N (unlikely)]			
		Were the treatment length, the therapeutic	regimen correct or in compliance with the SPC		
	1.4.				
		recommendation? [A clear NO points towar	•		
		Was the administration route used in comp	liance with the SPC recommendation? [A clear		
	1.5.				
		NO points towards N (unlikely)]			
	1.0	Was there a clear medicinal contra-indication	on for the products administered concurrently?		
	1.6.	TA also MEC as a later and ALC all al M			
		[A clear YES points towards N (unlikely)]			
	l '-	ng on the answers to questions 1.1. to 1.6, available conclude that the recommendations	an overall assessment of the information of the SPC :		
	o have	e been followed (all YES) e not been followed (one NO is enough) e difficult to conclude (unknown)	Points towards B (possible) or A (probable) Points directly towards N (unlikely) Points towards B (possible) or O1 (inconclusive) or O		
during	g the perio	of the clinical signs occur after the treatmen od of the efficacy of the product? [onset and n absence of specific clinical signs (presence	evolution of the clinical signs and/or presence of		
	Was the LEE identified during the efficacy period of the product? (if one efficacy period of the product is known)? [a clear NO points towards N (unlikely)]				
	YES		Points towards B (possible) or A (probable) Points		
	NO		directly towards N (unlikely) Points towards B (possible) or O1 (inconclusive) or O		
	It is not po	ossible to conclude	(unclassifiable/unassessable)		
3. Did th	ne clinical	signs fit the condition for which the product	is indicated?		
	3.1.		clinical signs of the adverse event recorded and PC? Are the clinical signs recorded specific or in line		





YES	Points towards B (possible) or A (probable) Points
NO It is r	directly towards N (unlikely) Points towards B (possible) or O1 (inconclusive) or O (unclassifiable/unassessable)



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4. Are there any measurable criteria to explain the event objectively? Has the diagnosis been confirmed? (post mortem results, laboratory results to confirm the diagnosis made before or after treatment of the animals or observations)? YES Points towards A (probable) NO/Unknown/Not applicable Points towards B (possible) or O1 (inconclusive) or O 5. Is there any information available concerning the farm environment that could explain the pathology (illness) despite animals having received treatment (if applicable)? 5.1. Was (were) the animal health status good? 5.2. Was the infestation pressure high? Is there any information related to concomitant pathology and the medical history of the breeding/ farming and/or of the animal? Are there reports of resistance to the product on 5.3. the farm or in the area where the event occurred? 5.4. Were zoo-technical and environmental measures taken? Were the hygiene conditions satisfactory? Were the farm management practices acceptable? An overview of the information allows conclusion that the environment factors Points towards B (possible) or O1 (inconclusive) or O Could explain in part (YES) (unclassifiable/unassessable) Did not play any role (NO) Points towards A (probable) It is not possible to conclude Points towards B (possible) or O1 (inconclusive) or O (unclassifiable/unassessable) 6. Is there any indication to confirm that the event is due to another cause that could explain the clinical signs recorded? There is a confirmed cause or aetiology indicating that the event is not due to/linked to Points directly towards N (unlikely) the non-efficacy of the product Points towards B (possible) or O1 (inconclusive) or O There are other plausible causes/explanations (unclassifiable/unassessable) There are no other causes/ explanations Points towards A (probable)





	A quality defect is suspected (e.g. storage conditions not respected)	Points towards B (possible) or O1 (inconclusive) or C (unclassifiable/unassessable)
	A quality defect is excluded (batch analysis available)	Points towards A (probable)
	No information available	Points towards B (possible) or O1 (inconclusive) or C (unclassifiable/unassessable) or A (probable)
	A quality defect has been clearly identified (batch analysis, expired batch)	This event should be assessed A (probable) This type of adverse event should be entered in EVVet but it should be clearly identified that the event is due to a batch quality defect. Indicate if the batch has been recalled.
8. Pr	evious knowledge of similar reports concerning the	LEE?
	8.1. There are scientific data	
	8.2.There are similar events reported	
	YES	Points towards B (possible) or A (probable)
	NO	Points towards B (possible) or O1 (inconclusive) or C (unclassifiable/unassessable) or A (probable)
Is the	reported information insufficient? Is there reason t	o doubt the reporting source/information?
	Yes	Points towards O1 (inconclusive) or O (unclassifiable/unassessable)