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SECTION 1: IDENTIFICATION		
1.1 Product identifier		
Product name:	Buprenodale Multidose®, 0.3mg/ml solution for injection Bupredine Multidose inj. 0,3 mg/ml 10 ml [ES/PT]	
Synonyms:	Not Available	
Proper Shipping name:	Not Available	
Other means of identification:	None	
1.2 Relevant identified uses	of the substances or mixture and uses advised against	
Recommended uses:	 Post-operative analgesia in the dog and cat. Post-operative analgesia, in combination with sedation, in the horse. Potentiation of the sedative effects of centrally acting agents in the dog and horse. 	
Uses advised against:	 Do not administer by the intrathecal or peridural route. The product should not be used pre-operatively in cases of Caesarean section, due to the risk of respiratory depression in the offspring periparturiently, and should only be used post-operatively with special care As reproductive studies have not been conducted in the target species, use only according to the benefit: risk assessment by the responsible veterinarian. Do not use in known cases of hypersensitivity to the active substance or any of the excipients. 	
1.3 Details of the supplier o	f the substance or mixture	
Registered company name:	Dechra Ltd	
Address:	Snaygill Industrial Estate Keighley Road Skipton North Yorkshire BD23 2RW UK	
Telephone:	+44 (0) 1756 791311	
Fax:	+44 (0) 1756 798604	
Email:	Not available	
1.4 Emergency Telephone N	Numbers	
	+44 (0) 1756 791311	

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SECTION 2: HAZARDS IDENTIFICATION		
2.1 Classification of the substance or mixture		
DSD Classification (EU):	Not Available	
DPD Classification (EU) ¹ :	Not Available	
Classification according to regulation (EC) No 1272/2008 [CLP] (EU) ¹ :	No data available	
2.2 Label Elements		
Signal Word:	No data available	
Hazard Statement(s)		
No data available		
Additional Statement(s)		
None		
Precautionary Statement(s)	Prevention:	
P270	Do not eat, drink or smoke when using this product.	
Precautionary Statement(s)	Response:	
P330	Rinse mouth.	
Precautionary Statement(s)	Precautionary Statement(s) Storage:	
P405	Store locked up	
Precautionary Statement(s) Disposal:		
P501	Dispose of contents / packaging according to local regulations	
2.3 Other Hazard Information N/a		

SECTION 3: INFORM	ATION ON THE I	NGREDIENTS	
3.1 Substances			
See section below for	composition of m	ixtures	
3.2 Mixtures			
1.CAS No 2.EC Number 3.Index Number 4.REACH Number	% Weight	Name	Classification according to regulations (EC) No 1272/2008 [CLP] (EU)

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1. 53152-21-9 2. 258-396-8 3. Not Available 4. Not Available	0.324%	Buprenorphine 0.3mg as buprenorphine hydrochloride 0.324mg	Acute Toxicity (Oral) Category 4, Reproductive Toxicity Category 2, Specific target organ toxicity - single exposure Category 3 (narcotic effects), Specific target organ toxicity - repeated exposure Category 2; H302, H361, H336, H373 [1]
N/a	Proprietary	Other ingredients determined not to be hazardous	N/a
Legend:	1. Classified by 0	Chemwatch	

SECTION 4: FIRST AID MEASURES		
4.1 Description of first a	aid measures	
Eye contact:	Following eye contamination, wash thoroughly with cold running water. Seek medical advice if irritation persists and show the package leaflet or the label to the medical practitioner.	
Skin contact:	Following skin contact, wash thoroughly with cold running water. Seek medical advice if irritation persists and show the package leaflet or the label to the medical practitioner.	
Inhalation:	Due to physical form of this product, inhalation exposure is unlikely. If accidentally inhaled, remove from exposure and seek medical attention if irritation occurs, showing the package leaflet or the label to the medical practitioner.	
Ingestion:	Remove material and flush mouth with water. Seek medical attention: treatment is symptomatic and includes replacement of fluid and electrolytes. Show the package leaflet or the label to the medical practitioner.	
Self-injection:	As buprenorphine has opioid-like activity, care should be taken to avoid accidental self-injection. In case of accidental self-injection or ingestion, seek medical advice immediately and show the package leaflet or the label to the medical practitioner.	
4.0. Most important symptoms and effects, both south and delayed		

4.2 Most important symptoms and effects, both acute and delayed

See Section 11

4.3 Indication of immediate medical attention and special treatment needed

Treat symptomatically.

Advice to doctors: Naloxone may be of benefit in reversing reduced respiratory rate and respiratory stimulants such as Doxapram are also effective in man. Because of the prolonged duration of effect of buprenorphine in comparison to such drugs, they may need to be administered repeatedly or by continuous infusion. Volunteer studies in man have indicated that opiate antagonists may not fully reverse the effects of buprenorphine.

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SECTION 5: FIRE FIGHTING MEASURES		
5.1 Extinguishing media		
Suitable:	Water spray, carbon dioxide, dry chemical and foam, as appropriate for surrounding fire and materials.	
Unsuitable:	None.	
5.2 Special hazards arising from the substance or mixture		
Fire incompatibility:	None known.	
5.3 Special protective actions for fire-fighters:		
Firefighting:	Alert Fire Brigade and tell them location and nature of hazard. Wear full breathing apparatus and self-contained breathing apparatus.	
Fire / explosion hazard:	None known.	

SECTION 6: ACCIDENTAL RELEASE MEASURES

6.1 Personal precautions, protective equipment and emergency procedures

For information on protective equipment, see section 8.

6.2 Environmental Precautions

Do not allow product to reach sewage system or any water course.

Inform respective authorities in case of seepage into water course or sewage system.

Do not allow to enter sewers/surface or ground water.

6.3 Methods and material for containment and cleaning up

6.5 Methods and material for containment and cleaning up		
Minor Spills:	Clean up all spills immediately. Avoid breathing vapours and contact with skin and eyes. Control personal contact with the substance, by using protective equipment. Contain and absorb spill with sand, earth, inert material or vermiculite.	
Major Spills:	Clear area of personnel and move upwind. Alert Fire Brigade and tell them location and nature of the hazard. Contain and absorb spill with sand, earth, inert material or vermiculite. Prevent, by any means available, spillage from entering drains or water course. Large spills should be collected into an appropriate container for waste disposal.	

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SECTION 7: HANDLING AND STORAGE			
7.1 Precautions for safe h	7.1 Precautions for safe handling		
Safe Handling:	Wear suitable protection gloves and clothing when handling the product. When handling, DO NOT eat, drink or smoke. Always wash hands with water after handling. Observe manufacturer's storage and handling recommendations.		
Other Information:	Protect from light. Keep out of the reach and sight of children.		
7.2 Conditions for safe sto	7.2 Conditions for safe storage, including any incompatibilities		
Suitable Container:	Shelf-life of the veterinary medicinal product as packaged for sale: 2 years. Shelf-life after first opening the vial: 28 days.		
Storage incompatibility:	Unknown.		
7.3 Specific end uses			
Not available			

SECTION 8: EXPOSURE CONTROLS / PERSONAL PROTECTION			
8.1 Control parameters	8.1 Control parameters		
DERIVED NO EFFECT LEVEL - DNEL (EU)			
Not Available	Not Available		
PREDICTED NO EFFECT LEVI	PREDICTED NO EFFECT LEVEL - PNEC (EU)		
Not Available	Not Available		
OCCUPATIONAL EXPOSURE LIMITS (OEL)			
INGREDIENT DATA:			
Not Available			
EMERGENCY LIMITS:			
Not Available			
8.2 Exposure controls			
	The basic types of engineering controls are: Process controls which involve changing the way a job activity or process is done to reduce the particular risk.		

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Personal protection:		
Eye and face protection:	Safety glasses with side shields / chemical goggles	
Skin protection:	See hand protection below	
Hands/ feet protection:	No special equipment needed when handling small quantities. OTHERWISE: Wear chemical protective gloves	
Body protection:	Wear appropriate clothing	
Other protection:	No special equipment needed when handling small quantities	
Thermal hazards:	Not applicable	
Respiratory protection:	Not applicable	
8.3 Environmental exposure controls See Section 12		

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SECTION 9: PHYSICAL AND CHEMICAL PROPERTIES

9.1 Information on basic physical and chemical properties

Appearance: Buprenodale Multidose®: Clear, colourless solution

Container: Cardboard carton containing a 10 ml clear Type I glass vial with a FluroTec

coated bromobutyl rubber stopper and aluminium seal.

Physical state: Solution for injection

Odour: Not available

Odour Threshold: Not available pH (as supplied): Not available

Melting point / freezing point (degrees C): Buprenorphine hydrochloride melting point:

265-271°C

Initial boiling point and boiling range: Not available

Flash Point: In water – no flash point. Evaporation rate: Not available Flammability: Not available

Upper/lower flammability or explosive limits: Not available

Vapour pressure: Not available

Relative Density (at degrees C): Not available

Solubility in water and solvents (mg/l): Buprenorphine hydrochloride in water: slightly

soluble

Vapour density: Not available

Auto ignition temperature (degrees C): Not available **Decomposition temperature (degrees C):** Not available

Viscosity: (degrees C): Not available Explosive properties: Not available Oxidising properties: Not available Partition Coefficient: Not available Molecular weight: Not available

Taste: Not available

Surface tension: Not available Volative component: Not available

Gas group: Not available pH as a solution: Not available

VOC g/L: Not available

9.2 Other information

Not Available

10: REACTIVITY AND STABILITY	
10.1 Reactivity:	See Section 7
10.2 Chemical stability:	Product is considered stable. Hazardous polymerisation will not occur.
10.3 Possibility of hazardous reactions:	The product is not considered to be hazardous if used as per instructions. Hazardous polymerisation will not occur.
10.4 Conditions to avoid:	Avoid excessive heat.

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10.5 Incompatible materials:	See section 7.
10.6 Hazardous decomposition:	See Section 5.

SECTION 11: TOXICOL	OGICAL INFORMATION		
Inhalation:	Due to physical form of this product, inhalation exposure is unlikely.		
Ingestion:	May cause irritation if ingested.		
Skin contact:	The product may produce sk	in irritation in some persons.	
Eye contact:	The product may produce eye irritation in some persons.		
Self-injection:	As buprenorphine has opioid-like activity, care should be taken to avoid accidental self-injection. In case of accidental self-injection or ingestion, seek medical advice immediately and show the package leaflet or the label to the physician.		
Chronic:	Naloxone may be of benefit in reversing reduced respiratory rate and respiratory stimulants such as Doxapram are also effective in man. Because of the prolonged duration of effect of buprenorphine in comparison to such drugs, they may need to be administered repeatedly or by continuous infusion. Volunteer studies in man have indicated that opiate antagonists may not fully reverse the effects of buprenorphine.		
Buprenodale Multidose®:	Acute toxicity	Irritation	
	Not Available	Not Available	
Buprenorphine hydrochloride:	Acute toxicity	Irritation	
	Oral (rat) LD ₅₀ : 80 mg/kg*e ¹	Eye: moderate Skin: moderate	
1.* Value obtained from manufacturer's SDS. Unless otherwise specified, data extracted from RTECS - Register of Toxic Effect of chemical Substances			
Skin corrosion/irritation	n:		
Not available			
Serious eye damage/ir	ritation:		
Not available			
Respiratory or skin sensitization:			
Not available			

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12.3 Bioaccumulative potential

No data Available

12.4 Mobility in SoilNo data Available

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SECTION 11: TOXICOLOGICAL INFORMATION
Germ cell mutagenicity:
Not available
Carcinogenicity:
Not available
Reproductive toxicity:
Laboratory studies in rats have not produced any evidence of a teratogenic effect. However, these studies have shown post-implantation losses and early foetal deaths. These may have resulted from a reduction in parental body condition during gestation and in post-natal care owing to sedation of the mothers.
STOT – single exposure:
Not available
STOT-repeated exposure:
Not available
Aspiration hazard:
Not available
SECTION 12: ECOLOGICAL INFORMATION
12.1 Toxicity
No data Available
DO NOT discharge into sewer or waterways.
12.2 Persistence and degradability
No data Available

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12.5 Results of PBT and vPvB assessment No data Available 12.6 Other adverse effects No data Available

SECTION 13: DISPOSAL CONSIDERATIONS		
13.1 Waste treatm	ent methods	
packaging	Any unused veterinary medicinal product or waste material derived from such veterinary medicinal products should be disposed of in accordance with national requirements.	
	Legislation addressing waste disposal requirements may differ by country, state and/ or territory. Each user must refer to laws operating in their area.	
	Ensure that the disposal of material is carried out in accordance with Hazardous Substances (Disposal) Regulations 2001.	
Waste Treatment Options:		
Sewage Disposal Options:	Not Available	

SECTION 14: TRANSPORT INFORMATION			
Labels required:	None		
Marine pollutant:	NO		
Hazchem:	Not Applicable		
Land transport (ADR):			
14.1 UN Numbe	er N/a	N/a	
14.2 UN Propo Shipping Nam			
14.3 Transport hazard class(es)		N/a	
	Sub risk	N/a	
14.4 Packing grou	p N/a	N/a	
14.5 Environment hazard			

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14.6 Special precautions for	Special provisions	N/a
user	Classification code	N/a
	Hazard Label	N/a
	Special provisions	N/a
	Limited quantity	N/a
14.7 Transport in bulk according to Annex II of MARPOL73/78 and the IBC Code		
Air transport (ICAO-l	ATA / DGR):	
14.1 UN Number	N/a	
14.2 UN Proper Shipping Name	N/a	
14.3 Transport	ICAO/IATA Class	N/a
hazard class(es)	ICAO / IATA Sub risk	N/a
	ERG Code	N/a
14.4 Packing group	N/a	
14.5 Environmental hazards	N/a	
	Special provisions	N/a
precautions for user	Cargo only packing instructions	N/a
	Cargo only maximum qty/pack	N/a
	Passenger and cargo packaging instructions	N/a
	Passenger and cargo maximum qty/pack	N/a
	Passenger and cargo limited quantity packing instructions	N/a
	Passenger and cargo limited maximum qty/pack	N/a

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14.7 Transport in bulk according to Annex II of MARPOL73/78 and the IBC Code	N/a			
Sea transport (IMDG	-Code / GGVSee):			
14.1 UN Number	N/a			
14.2 UN Proper Shipping Name	N/a			
14.3 Transport	IMDG Class	N/a		
hazard class(es)	IMDG Sub risk	N/a		
14.4 Packing group	N/a			
14.5 Environmental hazards	N/a			
	EMS Number	N/a		
precautions for user	Special provisions N/a			
	Limited quantities N/a			
14.7 Transport in bulk according to Annex II of MARPOL73/78 and the IBC Code	N/a			
Inland waterways tra	nsport (ADN):			
14.1 UN Number	r N/a			
14.2 UN Proper Shipping Name				
14.3 Transport hazard class(es)			N/a	
14.4 Packing group	N/a			
14.5 Environmental hazard	1			
	Classification Code		N/a	
precautions for user			N/a N/a	
	,			
	Special provisions		N/a	

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14.7 Transport in bulk according to Annex II of MARPOL73/78 and the IBC Code

SECTION 15: REGULATORY INFORMATION

15.1 Safety, health and environmental regulations / legislation specific for the substance or mixture

BUPRENORPHINE HYDROCHLORIDE (53152-21-9) IS FOUND ON THE FOLLOWING REGULATORY LISTS:

- European Customs Inventory of Chemical Substances ECICS (English)
- European Union European Inventory of Existing Commercial Chemical Substances (EINECS) (English)

This safety data sheet is in compliance with the following EU legislation and its adaptations - as far as applicable-: 98/24/EC, 92/85/EC, 94/33/EC, 91/689/EEC, 1999/13/EC

15.2 Chemical Safety Assessment

ECHA SUMMARY

Ingredient	CAS number	Index Number	ECHA Dossier
Buprenorphine hydrochloride	53152-21-9	Not Available	Not Available
Harmonization (C&L Inventory)	Hazard Class and Category Code(s)	Pictograms Signal Word Code(s)	Hazard Statement Code(s)
1	Acute Tox. 4, Repr. 2	GHS08, Wng	H302, H361
2	Acute Tox. 4, Repr. 2, Skin Sens. 1, Resp. Sens. 1, STOT SE 3, Lact., STOT SE 1, Acute Tox. 3, Acute Tox. 2, Repr. 1B, Eye Irrit. 2, STOT RE 2	GHS08, Dgr, GHS06	H317, H334, H336, H362, H370, H330, H300, H310, H360D, H319, H373

Harmonization Code 1 = The most prevalent classification. Harmonization Code 2 = The most severe classification

Australia - AICS	Υ
Canada - DSL	N (buprenorphine hydrochloride)
Canada - NDSL	N (buprenorphine hydrochloride)
China - IECSC	N (buprenorphine hydrochloride)
Europe - EINEC / ELINCS / NLP	Y

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Japan - ENCS	N (buprenorphine hydrochloride)
Korea - KECI	N (buprenorphine hydrochloride)
New Zealand - NZIoC	Υ
Philippines - PICCS	N (buprenorphine hydrochloride)
USA - TSCA	N (buprenorphine hydrochloride)
Legend:	Y = All ingredients are on the inventory N = Not determined or one or more ingredients are not on the inventory and are not exempt from listing(see specific ingredients in brackets)

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SECTION 16: OTHER INFORMATION

The SDS is written in accordance to guidelines specified by REACH, GHS and ECHA.

For detailed advice on Personal Protective Equipment, refer to the following EU CEN Standards:

EN 166 Personal eye-protection

EN 340 Protective clothing

EN 374 Protective gloves against chemicals and micro-organisms

EN 13832 Footwear protecting against chemicals

EN 133 Respiratory protective devices

Definitions and abbreviations

PC—TWA: Permissible Concentration-Time Weighted Average PC—STEL: Permissible Concentration-Short Term Exposure Limit

STEL: Short Term Exposure Limit

TEEL: Temporary Emergency Exposure Limit。

IDLH: Immediately Dangerous to Life or Health Concentrations

Version update Feb21: Updated product name to include Bupredine for ES/PT

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