This SDS packet was issued with item:

078935961

The safety data sheets (SDS) in this packet apply to the individual products listed below. Please refer to invoice for specific item number(s).

078934167 078934168 078935963 078935964 078937440 078937543 078939786 078939787 078939788 078939789 078939790 078939791



Gabapentin Capsules USP

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1. IDENTIFICATION

Material Gabapentin Capsules USP 100 mg, 300 mg and 400 mg

Recommended Use Rx Pharmaceutical for human use

Manufacturer Alkem Laboratories Ltd.

Mumbai - 400013, INDIA.

Distributor Ascend Laboratories, LLC

Parsippany, NJ 07054

Contact Phone Number 001-201-476-1977

2. HAZARD(S) IDENTIFICATION

GHS – Classification Not Classified as hazardous

Label Elements

Signal Word: Not classified

Hazard Statements: Not classified in accordance with international standards for

workplace safety

Other Hazards An Occupational Exposure Value has been established for one or

more of the ingredients (see Section 8).

Note This document has been prepared in accordance with standards for

workplace safety, which require the inclusion of all known hazards of the product or its ingredients regardless of the potential risk. The precautionary statements and warnings included may not apply in all cases. Your needs may vary depending upon the potential for

exposure in your workplace.

3. COMPOSITION / INFORMATION ON INGREDIENTS

Active Ingredient CAS No. Quantity

Gabapentin 60142-96-3 100 mg/Capsules, 300 mg/Capsules, and 400 mg/Capsules

Inactive IngredientCAS No.Inactive IngredientCAS No.Starch9005258Sodium Lauryl Sulfate151213



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Talc	14807966	Titanium Dioxide	13463677	
Lactose	63423 9000708	Ferric Oxide Yellow	51274001	
Gelatin	7000700	Ferric Oxide Red	1309371	

4. FIRST-AID MEASURE

Eye Contact Flush with water while holding eyelids open for at least 15 minutes.

Seek medical attention immediately.

Skin Contact Remove contaminated clothing. Flush area with large amounts of

water. Use soap. Seek medical attention.

Ingestion Never give anything by mouth to an unconscious person. Wash out

mouth with water. Do not induce vomiting unless directed by medical

personnel. Seek medical attention immediately.

Inhalation Remove to fresh air and keep patient at rest. Seek medical attention

immediately.

Most Important Symptoms and Effects, Both Acute and Delayed

Symptoms and For information on potential signs and symptoms of exposure, See

Effects of Exposure Section 2 – Hazards Identification and/or Section 11 - Toxicological

Information.
None known

Medical Conditions

Aggravated by

Exposure

Indication of the Immediate Medical Attention and Special Treatment Needed

Notes to Physician None

5. FIRE-FIGHTING MEASURE

Suitable Extinguishing Carbon dioxide, dry chemical or water.

Media

Special Hazards from Arising from the Substance or Mixture

Hazardous Combustion Formation of toxic gases is possible during heating or fire

Products

Fire/Explosion Hazards Fine particles (such as dust and mists) may fuel fires/explosions **Advice for Fire-Fighters** During all firefighting activities, wear appropriate protective



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equipment, including self-contained breathing apparatus

6. ACCIDENTAL RELEASE MEASURES

Personal Precautions

Protective Equipment and Emergency

Procedure

Personnel involved in clean-up should wear appropriate personal

protective equipment (see Section 8). Minimize exposure.

Environmental

Place waste in an appropriately labeled, sealed container for disposal.

Precautions Care should be taken to avoid environmental release.

Method and Material for Containment and Cleaning-up

Measures for

Cleaning/Collecting

Contain the source of spill if it is safe to do so. Collect spilled material by a method that controls dust generation. A damp cloth or a filtered

vacuum should be used to clean spills of dry solids. Clean spill area

thoroughly.

Additional

Consideration for

Large Spills

Non-essential personnel should be evacuated from affected area. Report emergency situations immediately. Cleanup operations should only be

undertaken by trained personnel.

7. HANDLING AND STORAGE

Precaution for Safe Handling Minimize dust generation and accumulation. If tablets or capsules are crushed and/or broken, avoid breathing dust and avoid contact with eyes, skin, and clothing. When handling, use appropriate personal protective equipment (see Section 8). Wash thoroughly after handling. Releases to the environment should be avoided. Review and implement appropriate technical and procedural waste water and waste disposal measures to prevent occupational exposure or environmental releases. Potential points of process emissions of this material to the atmosphere should be controlled with dust collectors, HEPA filtration systems or other equivalent controls.

Storage Conditions

Store at 20° to 25°C (68° to 77°F); [see USP Controlled Room

Temperature].

Specific End Use

Pharmaceutical drug product



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8. EXPOSURE CONTROLS / PERSONAL PROTECTION

Appropriate Engineering Controls

General room ventilation is adequate unless the process generates dust, mist or fumes. Engineering controls should be used as the primary means to control exposures. Keep airborne contamination levels below the exposure limits listed above in this section.

Personal Protective Equipment

Refer to applicable national standards and regulations in the selection and use of personal protective equipment (PPE). Contact your safety and health professional or safety equipment supplier for assistance in selecting the correct protective clothing/equipment based on an assessment of the workplace conditions, other chemicals used or present in the workplace and specific operational processes.

Hands

Impervious gloves (e.g. Nitrile, etc.) are recommended if skin contact with drug product is possible and for bulk processing operations. (Protective gloves must meet the standards in accordance with EN374, ASTM F1001 or international equivalent.)

Eyes

Wear safety glasses or goggles if eye contact is possible. (Eye protection must meet the standards in accordance with EN166, ANSI Z87.1 or international equivalent.)

Skin

Impervious protective clothing is recommended if skin contact with drug product is possible and for bulk processing operations. (Protective clothing must meet the standards in accordance with EN13982, ANSI 103 or international equivalent).

Respiratory Protection Under normal conditions of use, if the applicable Occupational Exposure Limit (OEL) is exceeded, wear an appropriate respirator with a protection factor sufficient to control exposures to below the OEL (e.g. particulate respirator with a half mask, P3 filter). (Respirators must meet the standards in accordance with EN140, EN143, ASTM F2704-10 or international equivalent).

9. PHYSICAL AND CHEMICAL PROPERTIES

Physical State Solid Form Capsules



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Sr. No.	Strength	Description	Packs	NDC No.
		White to off white crystalline	Bottles of 100	67877-222-01
		powder filled in size "4" white	Bottles of 500	67877-222-05
1	100 mg	hard gelatin capsule imprinted	Bottles of 1000	67877-222-10
		"216" on body with blue ink.	One Carton of 100	67877-222-38
		210 on body with blue lik.	(10 x 10 Unit-dose capsules)	0/8//-222-38
		White to off white crystalline	Bottles of 100	67877-223-01
	2 300 mg	powder filled in size "1"	Bottles of 500	67877-223-05
2		yellow colour hard gelatin	Bottles of 1000	67877-223-10
		capsule imprinted "215" on	One Carton of 100	67877-223-38
		body with blue ink.	(10 x 10 Unit-dose capsules)	0/8//-223-38
		White to off white crystalline	Bottles of 100	67877-224-01
		powder filled in size "0"	Bottles of 500	67877-224-05
3	400 mg	400 mg Orange colour hard gelatin	Bottles of 1000	67877-224-10
		capsule imprinted "214" on	One Carton of 100	67877-224-38
		bod y with blue ink.	(10 x 10 Unit-dose capsules)	0/0//-224-30

Explosive Properties Not explosive

Oxidizing Properties The substance or mixture is not classified as oxidizing

10. STABILITY AND REACTIVITY

Reactivity No data available.

Chemical Stability Stable at normal conditions of use.

Possibility of Hazardous Reactions

Polymerization Will not occur

Conditions to Avoid Fine particles (such as dust and mists) may fuel fires/explosions **Incompatible Materials** As a precautionary measure, keep away from strong oxidizers.

11. TOXICOLOGICAL INFORMATION

Information on Toxicological Effects

Short Term Dust may cause irritation (based on components). The active ingredient is

not acutely toxic.

Known Clinical Adverse effects associated with therapeutic use include dizziness, tiredness,



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Effects swelling, and nausea

Acute Toxicity (Species, Route, End Point, Dose):

Gabapentin

Species	Route	End Point	Dose
Mouse	Oral	LD50	> 5000 mg/kg
Rat	Oral	LD50	> 5000mg/kg
Rat	IV	LD50	> 2000mg/kg
Mouse	IV	LD50	1000-2000mg/kg
Rat	Subcutaneous	LD50	> 4000mg/kg

Talc (non-asbestiform)

Species	Route	End Point	<u>Dose</u>

Rat Oral LD50 > 1600 mg/kg

Acute Toxicity Comments:

A greater than symbol (>) indicates that the toxicity endpoint being tested was not achievable at the highest dose used in the test.

<u>Irritation / Sensitization (Study Type, Species, Severity):</u>

Gabapentin

Study TypeSpeciesSeverityEye IrritationRabbitNon-irritating

Repeated Dose Toxicity (Duration, Species, Route, Dose, End Point, Target Organ):

Gabapentin

Duration	Species	Route	Dose	End Point	Target Organ
52 weeks	Rat	Oral	250 mg/kg/day	NOAEL	Liver, Kidney
52 weeks	Monkey	Oral	250 mg/kg/day	NOAEL	Not identified
13 weeks	Mouse	Oral	1000 mg/kg/day	NOAEL	No effect at max dose

Reproduction & Developmental Toxicity (Study Type, Species, Route, Dose, End Point,

Effect(s)):

Gabapentin

Study Type	Species	Route	Dose	End Point	Effect(s)
Reproductive &	Rat	Oral	500	NOAEL	Negative
Fertility			mg/kg/day		
Embryo/Fetal	Mouse	Oral	3000	NOAEL	No effects at max dose
Development			mg/kg/day		
Embryo/Fetal	Rat	Oral	300	NOAEL	Developmental toxicity,



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Development mg/kg/day Not Teratogenic

Peri-/Postnatal Rat Oral 500 NOAEL Negative

Development mg/kg/day

Genetic Toxicity (Study Type, Cell Type/Organism, Result):

Gabapentin

Study Type	Cell Type/Organism,	Result
Bacterial Mutagenicity (Ames)	Salmonella, E. coli	Negative
In Vitro Chromosome Aberration	Hamster Lung Cells	Negative
In Vivo Unscheduled DNA Synthesis	Rat Hepatocyte	Negative
In Vivo Chromosome Aberration	Hamster Bone Marrow	Negative

Carcinogenicity (Duration, Species, Route, Dose, End Point, Effect(s)):

Gabapentin

Duration	Species	Route	Dose	End Point	Effect(s)
2 Year(s)	Mouse	Oral, in feed	2000 mg/kg/day	NOEL	Not carcinogenic
2 Year(s)	Male Rat	Oral, in feed	1000 mg/kg/day	NOEL	Malignant tumors,
					Pancreas

12. ECOLOGICAL INFORMATION

Environmental The environmental characteristics of this material have not been fully

Overview evaluated. Releases to the environment should be avoided.

Toxicity No data available **Persistence and** No data available

Degradability

Bio-accumulative Potential:

Gabapentin

Partition Coefficient:

MethodpHEndpointValuePredicted7.4Log D-1.31

Mobility in Soil: No data available

13. DISPOSAL CONSIDERATION

Waste Treatment Waste Treatment Methods: Dispose of waste in accordance with all applicable laws and regulations. Member State specific and

Community specific provisions must be considered. Considering the



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relevant known environmental and human health hazards of the material, review and implement appropriate technical and procedural waste water and waste disposal measures to prevent occupational exposure and environmental release. It is recommended that waste minimization be practiced. The best available technology should be utilized to prevent environmental releases. This may include destructive techniques for waste and wastewater.

14. TRANSPORT INFORMATION

The following refers to all modes of transportation unless specified below.

Not regulated for transport under USDOT, EUADR, IATA, or IMDG regulations.

15. REGULATORY INFORMATION

Generic Medicine. Approved by USFDA & the ANDA Number is 090858.

Safety, Health and Environmental Regulations/Legislation Specific for the Substance or **Mixture**

Gabapentin

Standard for the Uniform Scheduling for drugs and poisons	Schedule 4
EU EINECS/ELINCS List	262-076-3
Starch	
Inventory - United States TSCA - Sect. 8(b)	Present
Australia (AICS):	Present
REACH - Annex IV - Exemptions from the obligations of Register	Present
EU EINECS/ELINCS List	232-679-6
Lactose	
Inventory - United States TSCA - Sect. 8(b)	Present
Inventory - United States TSCA - Sect. 8(b) Australia (AICS)	Present Present
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Australia (AICS)	Present
Australia (AICS) REACH - Annex IV - Exemptions from the obligations of Register	Present Present
Australia (AICS) REACH - Annex IV - Exemptions from the obligations of Register EU EINECS/ELINCS List	Present Present
Australia (AICS) REACH - Annex IV - Exemptions from the obligations of Register EU EINECS/ELINCS List Talc	Present Present 200-559-2



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16. OTHER INFORMATION

The above information is based on data available to us and is believed to be correct. Since the information may be applied under conditions beyond our control and with which we may be unfamiliar, we do not assume any responsibility for the results of its use and all persons receiving it must make their own determination of the effects, properties and protections, which pertain to their particular conditions.

Alkem or Ascend shall not be held liable for any damage resulting from handling or from contact with the above product. Alkem or Ascend reserves the right to revise this SDS.

The information in this safety data sheet is based on current scientific knowledge. It should not be taken as expressing or implying any warranty concerning product characteristics.

Date of Preparation: 26/11/2019