1. IDENTIFICATION OF THE SUBSTANCE AND THE COMPANY

Material: Desloratadine Tablets, 5 mg

Manufacturer: Lupin Limited
Goa 403 722
INDIA

Distributor: Lupin Pharmaceuticals, Inc.
Harborplace Tower, 21st Floor
111, South Calvert Street
Baltimore, MD 21202
United States
Tel. 001-410-576-2000
Fax. 001-410-576-2221

2. COMPOSITION / INFORMATION ON INGREDIENTS

<table>
<thead>
<tr>
<th>Ingredients</th>
<th>CAS</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Desloratadine</td>
<td>100643-71-8</td>
<td>5 mg</td>
</tr>
</tbody>
</table>

3. HAZARD IDENTIFICATION

Fire and Explosion: Expected to be non-combustible

Health: Desloratadine tablets are contraindicated in patients who are hypersensitive to this medication or to any of its ingredients or to loratadine.

Environment: No information is available about the potential of this product to produce adverse environmental effects.

4. FIRST AID MEASURE

Ingestion: If conscious, give water to drink and induce vomiting. Do not attempt to give any solid or liquid by mouth if the exposed subject is unconscious or semi-conscious. Wash out the mouth with water. Obtain medical attention.
Inhalation
Move individual to fresh air. Obtain medical attention if breathing difficulty occurs. If not breathing, provide artificial respiration assistance.

Skin Contact
Remove contaminated clothing and flush exposed area with large amounts of water. Wash all exposed areas of skin with plenty of soap and water. Obtain medical attention if skin reaction occurs.

Eye Contact
Flush eyes with plenty of water. Get medical attention.

NOTES TO HEALTH PROFESSIONALS

Medical Treatment
Treat according to locally accepted protocols. For additional guidance, refer to the current prescribing information or to the local poison control information center. Protect the patient’s airway and support ventilation and perfusion. Meticulously monitor and maintain, within acceptable limits, the patient’s vital signs, blood gases, serum electrolytes, etc.

Overdosage
In the event of overdose, consider standard measures to remove any unabsorbed drug. Symptomatic and supportive treatment is recommended. Desloratadine and 3-hydroxydesloratadine are not eliminated by hemodialysis.

Information regarding acute overdosage is limited to experience from post-marketing adverse event reports and from clinical trials conducted during the development of the desloratadine product. In a dose-ranging trial, at doses of 10 mg and 20 mg/day somnolence was reported.

In another study, no clinically relevant adverse events were reported in normal male and female volunteers who were given single daily doses of desloratadine 45 mg for 10 days.

5. FIRE FIGHTING MEASURE

Fire and Explosion Hazards
Assume that this product is capable of sustaining combustion.

Extinguishing Media
Water spray, carbon dioxide, dry chemical powder or appropriate foam.

Special Firefighting Procedures
For single units (packages): No special requirements needed. For larger amounts (multiple packages/pallets) of product: Since toxic, corrosive or flammable vapors might be evolved from fires involving this product and associated packaging, self contained breathing apparatus and full protective equipment are recommended for firefighters.

Hazardous Combustion Products
Hazardous combustion or decomposition products are expected when the product is exposed to fire.
6. ACCIDENTAL RELEASE MEASURES

Personal Precautions
Wear protective clothing and equipment consistent with the degree of hazard.

Environmental Precautions
For large spills, take precautions to prevent entry into waterways, sewers, or surface drainage systems.

Clean-up Methods
Collect and place it in a suitable, properly labeled container for recovery or disposal.

7. HANDLING AND STORAGE

Handling
No special control measures required for the normal handling of this product.

Heat Sensitive.
Avoid exposure at or above 30°C (86°F).
Dispense in tight, light-resistant container as defined in the USP using a child-resistant closure.

Storage
Store at 25°C (77°F); excursions permitted to 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature].

8. EXPOSURE CONTROLS / PERSONAL PROTECTION

Wear appropriate clothing to avoid skin contact. Wash hands and arms thoroughly after handling.

9. PHYSICAL AND CHEMICAL PROPERTIES

Physical Form
Desloratadine Tablets, 5 mg are light blue, circular, biconvex, film-coated tablets debossed "LU" on one side and "S71" on other side.
They are supplied as follows:

<table>
<thead>
<tr>
<th>NDC</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>68180-153-01</td>
<td>Bottles of 100</td>
</tr>
<tr>
<td>68180-153-02</td>
<td>Bottles of 500</td>
</tr>
<tr>
<td>68180-153-12</td>
<td>3x10’s unit dose blisters</td>
</tr>
<tr>
<td>68180-153-13</td>
<td>10x10’s unit dose blisters</td>
</tr>
</tbody>
</table>

10. STABILITY AND REACTIVITY

Stable under recommended storage conditions.
11. TOXICOLOGICAL INFORMATION

Carcinogenicity Studies

The carcinogenic potential of desloratadine was assessed using a loratadine study in rats and a desloratadine study in mice. In a 2-year study in rats, loratadine was administered in the diet at doses up to 25 mg/kg/day (estimated desloratadine and desloratadine metabolite exposures were approximately 30 times the AUC in humans at the recommended daily oral dose). A significantly higher incidence of hepatocellular tumors (combined adenomas and carcinomas) was observed in males given 10 mg/kg/day of loratadine and in males and females given 25 mg/kg/day of loratadine. The estimated desloratadine and desloratadine metabolite exposures in rats given 10 mg/kg of loratadine were approximately 7 times the AUC in humans at the recommended daily oral dose. The clinical significance of these findings during long-term use of desloratadine is not known.

In a 2-year dietary study in mice, males and females given up to 16 mg/kg/day and 32 mg/kg/day desloratadine, respectively, did not show significant increases in the incidence of any tumors. The estimated desloratadine and desloratadine metabolite exposures in mice at these doses were 12 and 27 times, respectively, the AUC in humans at the recommended daily oral dose.

Genotoxicity Studies

In genotoxicity studies with desloratadine, there was no evidence of genotoxic potential in a reverse mutation assay (Salmonella/E. coli mammalian microsome bacterial mutagenicity assay) or in 2 assays for chromosomal aberrations (human peripheral blood lymphocyte clastogenicity assay and mouse bone marrow micronucleus assay).

Impairment of Fertility

There was no effect on female fertility in rats at desloratadine doses up to 24 mg/kg/day (estimated desloratadine and desloratadine metabolite exposures were approximately 130 times the AUC in humans at the recommended daily oral dose). A male specific decrease in fertility, demonstrated by reduced female conception rates, decreased sperm numbers and motility, and histopathologic testicular changes, occurred at an oral desloratadine dose of 12 mg/kg in rats (estimated desloratadine and desloratadine metabolite exposures were approximately 45 times the AUC in humans at the recommended daily oral dose). Desloratadine had no effect on fertility in rats at an oral dose of 3 mg/kg/day (estimated desloratadine and desloratadine metabolite exposures were approximately 8 times the AUC in humans at the recommended daily oral dose).

12. ECOLOGICAL INFORMATION

No relevant studies identified.
13. DISPOSAL CONSIDERATION

Incinerate in an approved facility. Follow all federal state and local environmental regulations.

14. TRANSPORT INFORMATION

<table>
<thead>
<tr>
<th>IATA/ICAO - Not Regulated</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>IATA Proper shipping Name</td>
<td>N/A</td>
</tr>
<tr>
<td>IATA UN/ID No</td>
<td>N/A</td>
</tr>
<tr>
<td>IATA Hazard Class</td>
<td>N/A</td>
</tr>
<tr>
<td>IATA Packaging Group</td>
<td>N/A</td>
</tr>
<tr>
<td>IATA Label</td>
<td>N/A</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>IMDG - Not Regulated</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>IMDG Proper shipping Name</td>
<td>N/A</td>
</tr>
<tr>
<td>IMDG UN/ID No</td>
<td>N/A</td>
</tr>
<tr>
<td>IMDG Hazard Class</td>
<td>N/A</td>
</tr>
<tr>
<td>IMDG Flash Point</td>
<td>N/A</td>
</tr>
<tr>
<td>IMDG Label</td>
<td>N/A</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>DOT - Not Regulated</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>DOT Proper shipping Name</td>
<td>N/A</td>
</tr>
<tr>
<td>DOT UN/ID No</td>
<td>N/A</td>
</tr>
<tr>
<td>DOT Hazard Class</td>
<td>N/A</td>
</tr>
<tr>
<td>DOT Flash Point</td>
<td>N/A</td>
</tr>
<tr>
<td>DOT Packing Group</td>
<td>N/A</td>
</tr>
<tr>
<td>DOT Label</td>
<td>N/A</td>
</tr>
</tbody>
</table>

15. REGULATORY INFORMATION

This Section Contains Information relevant to compliance with other Federal and/or state laws.

16. OTHER INFORMATION

The above information is believed to be correct but does not purport to be all-inclusive and shall be used only as a guide. Nothing herein shall be deemed to create any warranty, express or implied. It is the responsibility of the user to determine the applicability of this information and the suitability of the material or product for any particular purpose.

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