# **Health Products Regulatory Authority**

# **Summary of Product Characteristics**

#### **1 NAME OF THE MEDICINAL PRODUCT**

Dymista 137 micrograms/50 micrograms per actuation nasal spray, suspension

## **2 QUALITATIVE AND QUANTITATIVE COMPOSITION**

Each g of suspension contains 1000 micrograms azelastine hydrochloride and 365 micrograms fluticasone propionate. Each actuation (0.14 g) delivers 137 micrograms azelastine hydrochloride (equivalent to 125 micrograms azelastine) and 50 micrograms fluticasone propionate.

Excipient with known effect: Benzalkonium chloride

For the full list of excipients, see section 6.1.

#### **3 PHARMACEUTICAL FORM**

**Nasal Spray Suspension** 

Product imported from France White, homogenous suspension.

#### **4 CLINICAL PARTICULARS**

As per PA23355/010/001

## **5 PHARMACOLOGICAL PROPERTIES**

As per PA23355/010/001

## **6 PHARMACEUTICAL PARTICULARS**

## 6.1 List of excipients

Disodium edetate Glycerol Microcrystalline cellulose Carmellose sodium Polysorbate 80 Benzalkonium chloride Phenylethyl alcohol Purified water

#### 6.2 Incompatibilities

Not applicable

#### 6.3 Shelf life

The shelf life expiry date of this product is the date shown on the bottle and outer carton of the product as marketed in the country of origin.

In-use shelf life (after first use): 6 months

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# **6.4 Special precautions for storage**

Do not refrigerate or freeze.

## 6.5 Nature and contents of container

Cardboard outer carton containing an amber coloured glass bottle fitted with a spray pump, applicator and a protective cap.

Pack size: 1 bottle with 23 g suspension in 25 ml bottles (at least 120 actuations)

# 6.6 Special precautions for disposal and other handling

No special requirements for disposal.

## **7 PARALLEL PRODUCT AUTHORISATION HOLDER**

IMED Healthcare Ltd. Unit 625 Kilshane Avenue Northwest Business Park Ballycoolin Dublin 15 Ireland

## **8 PARALLEL PRODUCT AUTHORISATION NUMBER**

PPA1463/151/001

# 9 DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

Date of first authorisation: 27<sup>th</sup> October 2017

#### 10 DATE OF REVISION OF THE TEXT

September 2023

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