

REC 0/0559 - Emerging Insight and Market Forecast - 2030

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Abstracts

This report can be delivered to the clients within 48 Hours

“REC 0/0559 - Emerging Insight and Market Forecast - 2030” the report provides comprehensive insights about an investigational product for Persistent Epithelial Defect in 7 Major Markets. A detailed picture of the REC 0/0559 in Seven Major Markets, i.e., United States, EU5 (Germany, France, Italy, Spain, and the United Kingdom), and Japan, for the study period 2020–2030 is provided in this report along with a detailed description of the product. The product details cover mechanism of action, dosage and administration, route of synthesis, and Research and development activity including regulatory milestones, and other development activities. Further, it also consists of future market assessments inclusive of the market forecast, SWOT analysis, market competitors, and other emerging therapies.

Overview

REC 0/0559 (also known as MT-8 or REC-0559) is a low molecular weight non-peptidic human nerve growth factor (NGF) mimetic currently under global development by Recordati. REC 0559 was licensed in 2017 from MimeTech, an Italian based company focused on the development of pharmaceutical applications for synthetic neurotrophin mimetics. FDA has granted Orphan Drug Designation to Recordati Rare Disease's investigational product REC 0559 for the treatment of neurotrophic keratitis.

Scope of the report

The report provides insights into:

A comprehensive product overview including the product description, mechanism of action, dosage and administration, Research and Development activity.

Elaborated details on regulatory milestones and other development activities have been provided in this report.

The report also highlights the drug research and development activity details across the United States, Europe and Japan.

The report also covers the patents information with expiry timeline around REC 0/0559.

The report contains forecasted sales for REC 0/0559 till 2030.

Comprehensive coverage of the late-stage emerging therapies (Phase II) for Persistent Epithelial Defect.

The report also features the SWOT analysis with analyst insights and key findings of REC 0/0559.

Methodology

The report is built using data and information sourced primarily from internal databases, primary and secondary research and in-house analysis by DelveInsight's team of industry experts. Information and data from the secondary sources have been obtained from various printable and nonprintable sources like search engines, news websites, global regulatory authorities websites, trade journals, white papers, magazines, books, trade associations, industry associations, industry portals and access to available databases.

REC 0/0559 Analytical Perspective by DelveInsight

In-depth REC 0/0559 Market Assessment

This report provides a detailed market assessment of REC 0/0559 in Seven Major Markets, i.e., United States, EU5 (Germany, France, Italy, Spain, and the United

Kingdom), and Japan. This segment of the report provides forecasted sales data from 2020 to 2030.

REC 0/0559 Clinical Assessment

The report provides the clinical trials information of REC 0/0559 covering trial interventions, trial conditions, trial status, start and completion dates.

Report highlights

In the coming years, the market scenario for Persistent Epithelial Defect is set to change due to the extensive research and incremental healthcare spending across the world; which would expand the size of the market to enable the drug manufacturers to penetrate more into the market.

The companies and academics are working to assess challenges and seek opportunities that could influence REC 0/0559 dominance. The therapies under development are focused on novel approaches to treat/improve the disease condition.

Other emerging products for Persistent Epithelial Defect are giving market competition to REC 0/0559 and launch of late-stage emerging therapies in the near future will significantly impact the market.

A detailed description of regulatory milestones, development activities, and some key findings provide the current development scenario of REC 0/0559.

Our in-depth analysis of the forecasted sales data of REC 0/0559 from 2020 to 2030 will support the clients in the decision-making process regarding their therapeutic portfolio by identifying the overall scenario of the REC 0/0559.

Key Questions

Which company is developing REC 0/0559 along with the phase of the clinical study?

What is the technology utilized in the development of REC 0/0559?

What is the product type, route of administration and mechanism of action of REC 0/0559?

What is the clinical trial status of the study and study completion date?

What are the key collaborations, mergers and acquisitions, licensing and other activities related to the REC 0/0559 development?

What are the key designations that have been granted to REC 0/0559?

What is the forecasted market scenario of REC 0/0559?

What is the history of REC 0/0559 and what is its future?

What is the forecasted sales of REC 0/0559 in the seven major countries, including the United States, Europe (Germany, France, Italy, Spain, and the United Kingdom), and Japan?

What are the other emerging products available and how these are giving competition to REC 0/0559?

Which are the late-stage emerging therapies under development for the treatment of the PPD?

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