

Esbriet (pirfenidone) - Drug Insight and Market Forecast - 2030

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Abstracts

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OVERVIEW

"Esbriet (pirfenidone) - Drug Insight and Market Forecast – 2030" report by DelveInsight outlays comprehensive insights of the product indicated for the treatment of its approved condition. A detailed picture of the Esbriet (pirfenidone) in Seven Major Markets, i.e., United States, EU5 (Germany, France, Italy, Spain, and the United Kingdom), and Japan, for the study period 2017–2030 is provided in this report along with a detailed description of the product. The product details covers mechanism of action, dosage and administration, route of synthesis, and pharmacological studies, including product marketed details, regulatory milestones, and other development activities. Further, it also consists of market assessments inclusive of the market forecast, SWOT analysis, and detailed analyst views. It further highlights the market competitors, late-stage emerging therapies, and patent details in the global space.

Esbriet is an oral anti-fibrotic medicine approved for the treatment of Idiopathic Pulmonary Fibrosis (IPF), and is available in approximately 40 countries worldwide.In clinical studies pirfenidone was better than placebo in improving lung function; however, it did not improve survival in people with idiopathic pulmonary fibrosis. ESBRIET is available as a white to off-white hard gelatin capsule containing 267 mg of pirfenidone for oral administration, or, as film-coated tablets containing 267 mg (yellow) and 801 mg (brown) pirfenidone. Esbriet has Orphan Drug Designation and was approved for use in Europe in 2011 in adults with mild-to-moderate IPF, and in the US in people with IPF in October 2014. In 2017, the U.S. Food and Drug Administration (FDA) and the European Commission approved the Esbriet 801 mg and 267 mg tablets as an alternative to the



original capsule formulation. The new 801 mg tablets offer people with IPF a maintenance option for taking Esbriet with fewer pills per day. In 2019, the U.S. Food and Drug Administration (FDA) granted Breakthrough Designation status for Esbriet in unclassifiable interstitial lung disease (uILD).

SCOPE OF THE REPORT

The report provides insights into:

A comprehensive product overview including the product description, mechanism of action, dosage and administration, route of synthesis, pharmacological studies (pharmacodynamics and pharmacokinetics) and adverse reactions.

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

The report also highlights the drug marketed details across the United States, Europe and Japan.

The report also covers the patents information with expiry timeline around Esbriet (pirfenidone).

The report contains historical and forecasted sales for Esbriet (pirfenidone) till 2030.

Comprehensive coverage of the late-stage emerging therapies (Phase III) in the space with a brief snapshot of the details.

The report also features the SWOT analysis with analyst insights and key findings of Esbriet (pirfenidone).

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.

Esbriet (pirfenidone) Analytical Perspective by DelveInsight

In-depth Esbriet (pirfenidone) Market Assessment

This report provides a detailed market assessment of Esbriet (pirfenidone) 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 historical and forecasted sales data from 2017 to 2030.

Esbriet (pirfenidone) Clinical Assessment

The report provides the clinical trials information of Esbriet (pirfenidone) covering trial interventions, trial conditions, trial status, start and completion dates.

REPORT HIGHLIGHTS

In the coming years, the market scenario for Esbriet (pirfenidone) is set to change due to the extensive research in the treatment of the indicated condition 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 Esbriet (pirfenidone) dominance. The therapies under development are focused on novel approaches to treat/improve the disease condition.

Other approved products for the disease are giving market competition to Esbriet (pirfenidone) 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 market scenario of Esbriet (pirfenidone).



Our in-depth analysis of the sales data of Esbriet (pirfenidone) from 2017 to 2030 will support the clients in the decision-making process regarding their therapeutic portfolio by identifying the overall scenario of the Esbriet (pirfenidone) in the market.

KEY QUESTIONS

What is the prescribed dosage and strengths of Esbriet (pirfenidone) are available in the market?

What are the common adverse reactions or side effects of Esbriet (pirfenidone)?

What is the product type, route of administration and mechanism of action of Esbriet (pirfenidone)?

What are the chemical specifications of Esbriet (pirfenidone)?

How are the clinical trials diversified on the basis of the trial status?

What is the history of Esbriet (pirfenidone), and what is its future?

What are the marketed details of Esbriet (pirfenidone) in the seven major countries, including the United States, Europe (Germany, France, Italy, Spain, and the United Kingdom), and Japan?

How many patents have been granted to Esbriet (pirfenidone) and when these patents will get expire?

What are the pros (benefits) and cons (disadvantages) of Esbriet (pirfenidone)?

In which countries Esbriet (pirfenidone) got approval and when it gets launched?

What are the clinical trials are currently ongoing for Esbriet (pirfenidone)?

How the safety and efficacy results determined the approval of Esbriet (pirfenidone)?



What are the key collaborations, mergers and acquisitions, licensing and other activities related to the Esbriet (pirfenidone) development?

What are the key designations that have been granted to Esbriet (pirfenidone)?

What is the historical and forecasted market scenario of Esbriet (pirfenidone)?

How is the market trend of Esbriet (pirfenidone) is different in the Seven Major Markets (the United States, EU5 [Germany, France, Italy, Spain, and the United Kingdom], and Japan)?

What are the other approved products available and how these are giving competition to Esbriet (pirfenidone)?

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



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