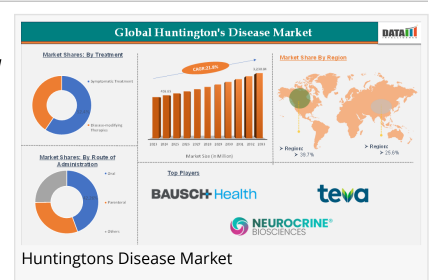


## Huntington's Disease Market to Reach \$3.2Bn by 2033 Amid Breakthroughs in Gene-Based Therapies | DataM Intelligence

Huntington's Disease market is set to hit \$3.2Bn by 2033, propelled by gene therapy innovation, rising awareness, and strong pharma R&D focus.

NEW YORK, NY, UNITED STATES, June 13, 2025 /EINPresswire.com/ -- Market Overview:

The <u>Huntington's Disease Market</u> is experiencing a surge in innovation and investment as pharmaceutical companies and researchers work



toward developing more effective and personalized treatments. Huntington's disease is a rare genetic disorder marked by worsening motor control, cognitive impairment, and behavioral or psychiatric symptoms. Due to the complex pathology of the disease, treatment options have traditionally been limited. However, increased understanding of the genetic mechanisms behind

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Innovative gene therapies and a rising focus on rare neurological disorders are reshaping the Huntington's Disease market and giving hope to patients worldwide." HD is spurring the development of advanced therapies, from gene silencing techniques to symptomatic relief medications. The market was valued at US\$ 455.65 million in 2024 and is projected to climb to US\$ 3,230.84 million by 2033, expanding at a 7.5% CAGR from 2025 to 2033.

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DataM Intelligence

Market Drivers are :

Growing prevalence of rare neurological disorders: Rising awareness and improved diagnosis of HD globally.

Increased R&D investments in neurodegenerative diseases: Major pharmaceutical firms are prioritizing orphan disease pipelines.

Advances in gene therapy and RNA interference (RNAi): Promising avenues for modifying disease progression.

Supportive regulatory frameworks: Accelerated approvals and orphan drug designations incentivize development.

Expanding access to healthcare and genetic testing: Early diagnosis facilitates timely therapeutic interventions.

Market Key Players are :

Teva Pharmaceutical Industries Ltd.

Bausch Health Companies Inc.

Neurocrine Biosciences, Inc.

Hetero

Lupin

Hikma Pharmaceuticals PLC

Dr. Reddy's Laboratories Ltd.

Sun Pharmaceutical Industries Ltd.

Novartis AG

uniQure NV

Market Segmentation:

By Drug Type: Antipsychotic Drugs, Antidepressants, Tetrabenazine, Anticonvulsants, Others

By Treatment Type: Symptomatic Treatment, Disease-Modifying Treatment

By Distribution Channel: , Retail Pharmacies, Online Pharmacies and Hospital Pharmacies

By Region: Europe, Asia-Pacific, South America, North America, Middle East & Africa

Latest News of USA:-

In 2024, Neurocrine Biosciences expanded clinical trials for valbenazine, targeting tardive dyskinesia and chorea in HD patients. The company also announced new partnerships with U.S. research institutions to accelerate gene therapy studies focused on silencing mutant huntingtin protein expression.

Latest News of Japan:-

Japan's Ministry of Health, Labour and Welfare approved expanded coverage for genetic testing of Huntington's disease, aiming to promote early diagnosis and family counseling. Additionally, collaborative research between Tokyo University and domestic biotech firms received government funding to explore RNA-targeted therapies for HD.

Recent Key Developments are :

Teva Pharmaceuticals launched a new educational program in North America to improve understanding of HD treatments among healthcare providers.

Bausch Health progressed with Phase 2 clinical trials of a novel small molecule intended to delay neurodegeneration in HD patients.

Neurocrine Biosciences secured FDA fast-track designation for its next-gen therapy aimed at HDassociated chorea.

## Conclusion:

The Huntington's Disease market is on a transformative trajectory, driven by growing awareness, scientific breakthroughs, and active engagement from key pharmaceutical companies. As clinical research continues to advance, particularly in disease-modifying and genetic therapies, the outlook for HD patients is gradually improving. With regulatory support and increased investment, the next decade could see significant progress in turning this devastating condition into a more manageable one.

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