

Asia-Pacific Rare Disease Genetic Testing Market: Analysis and Forecast, 2023-2033

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Abstracts

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Introduction to Asia-Pacific Rare Disease Genetic Testing Market

The Asia-Pacific rare disease genetic testing market was valued at \$0.58 billion in 2023 and is anticipated to reach \$2.40 billion by 2033, witnessing a CAGR of 15.25% during the forecast period 2023-2033. Advanced technologies like whole-genome sequencing (WGS) and whole-exome sequencing (WES) analyze an individual's entire genetic code (genome) or specific protein-coding regions (exome), respectively. They aid in detecting genetic variations, mutations, or abnormalities linked to rare diseases.

Market Introduction

The Asia-Pacific (APAC) rare disease genetic testing market is witnessing notable growth driven by various factors. Increased awareness, advancements in genetic testing technologies, and rising healthcare expenditure contribute to market expansion. Moreover, the region's large population base and diverse genetic profiles present significant opportunities for genetic testing providers. Government initiatives aimed at improving healthcare infrastructure and expanding access to genetic testing services further propel market growth. Additionally, collaborations between healthcare organizations, research institutions, and industry players facilitate the development and adoption of innovative genetic testing solutions tailored to the region's specific needs. With a growing focus on precision medicine and personalized healthcare, the APAC rare disease genetic testing market is poised for continued expansion and innovation in



the foreseeable future.

Market Segmentation:

Segmentation 1: by Disease Type

Gastroenterology Disease

Endocrine and Metabolism Disease

Cardiovascular Disease

Neurology Disease

Hematology and Oncology Disease

Dermatology Disease

Other Diseases

Segmentation 2: by Technology

Sanger Sequencing

PCR

NGS

Microarrays

Others (Targeted Cytogenetics, Karyotyping, and FISH)

Segmentation 3: by Country

Japan

India



China

South Korea

Australia

Singapore

Rest-of-Asia-Pacific

How Can This Report Add Value to an Organization?

Product/Innovation Strategy: The APAC rare disease genetic testing market has been extensively segmented based on various categories, such as disease type, technology, and country. This can help readers understand which segments account for the largest share and which are well-positioned to grow in the coming years.

Competitive Strategy: The APAC rare disease genetic testing market has numerous established players paving their way into providing tests, panels, and assays in the market. Key players in the global rare disease genetic testing market analyzed and profiled in the study involve established players offering various disease-specific panels and multiplex instruments.

Key Market Players and Competition Synopsis

The companies that are profiled have been selected based on inputs gathered from primary experts and analysing the company's coverage, product portfolio, its market penetration.

Some of the prominent names established in this market are:

3billion, Inc.

Konica Minolta (Ambry Genetics, Inc.)

BGI Group



Strand Life Science Pvt Ltd.



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