

Neurodegenerative Diseases (AD, PD, HD) Diagnostics

1. Introduction

What are Neurodegenerative Diseases?

Neurodegenerative diseases are defined as diseases described by the progressive dysfunction or death of neurons or nerve cells in the human body. These diseases mainly target the central nervous system meaning the function of the affected body gradually diminishes and reaches a state of severe disability or the patient dies. Some of the familiar brain disorders associated with neurodegenerative diseases are Parkinson's Disease, Alzheimer's Disease and Huntington's Disease ^[1-3].

The main characteristics of these diseases are:

- Progressive Deterioration: Symptoms and functions worsen over time.
- **Irreversibility:** Currently, most neurodegenerative diseases have no cure, and treatments can only alleviate symptoms and slow disease progression.
- **Multifactorial Etiology:** These diseases are influenced by both genetic and environmental as well as lifestyle factors.

Why is early diagnosis important?

For several reasons, neurodegenerative disease requires early diagnosis and effective treatment.

- **Improving Quality of Life:** Early detection and intervention can slow disease progression, alleviate symptoms, and enhance the quality of life for patients.
- **Prolonging Life:** Early treatment can extend the life expectancy of patients by delaying the severe outcomes associated with these diseases.
- **Reducing Societal Burden:** Early diagnosis and treatment can lower healthcare costs, reduce the economic and psychological burden on patients and their families, and lessen the overall societal impact.
- **Promoting Scientific Research:** More cases and data with early diagnosis means more to study disease mechanisms, and hopefully learn how to develop new therapies.

2. Common Degenerative Diseases

Neurodegenerative diseases are a group of disorders characterized by the gradual degeneration or death of neurons, the fundamental units of the nervous system. These conditions primarily affect the central nervous system, leading to a progressive loss of function, which can ultimately result in severe disability or death. Common neurodegenerative diseases include Parkinson's disease, Alzheimer's disease, and Huntington's disease.



Alzheimer's Disease (AD)

The most common form of dementia, Alzheimer's disease is a neurodegenerative disorder largely defined by the memory loss and subsequent drop in cognitive ability. The disease is almost always of the elderly, predominantly over the age of 65. But some people get early onset Alzheimer's disease, and start to exhibit symptoms earlier than that ^[4,5].

Pathological Changes

The neuropathological hallmarks of Alzheimer's disease primarily include:

- **Amyloid Plaques:** These are aggregates formed from protein fragments due to the abnormal cleavage of amyloid precursor protein.
- **Neurofibrillary Tangles:** These tangles occur because abnormal tau protein builds up in the cytoplasm of neurons, and disrupts the intracellular transport system.

Diagnosis

- 1. Clinical Symptom Assessment
- 2. Neuropsychological Testing
- 3. Brain Imaging Techniques
- 4. Biomarker Testing (Aβ42, p-tau217, p-tau181, p-tau231, GFAP, NfL)

Parkinson's Disease (PD)

Parkinson's disease is a neurodegenerative disorder that primarily affects the motor system. Such a loss of dopamine producing neurons in the substantia nigra region of the brain causes the disease, which results in less dopamine. Dopamine is as important a neurotransmitter as it is to the movement as it is to the emotional responses. These neurons are damaged or killed leading to the characteristic motor and non-motor symptoms seen in Parkinson's disease ^[6,7].

Pathological Changes

- 1. Loss of Dopamine Neurons
- 2. Formation of Lewy Bodies
- 3. Neurotransmitter Changes
- 4. Neurotransmitter Changes



Diagnosis

- 1. Clinical Symptom Assessment
- 2. Neurological Examination
- 3. Drug Response Test
- 4. Imaging Studies
- 5. Biomarker Testing (Alpha-synuclein, DJ-1, NfL)

Huntington's Disease (HD)

A CAG repeats expansion mutation of the HTT gene leading to the abnormal accumulation of the mutant huntingtin protein (mHTT) is a hereditary rare neurodegenerative disorder ^[8,9].

Pathological Changes

- 1. Neuronal Loss
- 2. Accumulation of Mutant Huntingtin Protein
- 3. Formation of Neuronal Inclusions
- 4. Impairment of Neural Transmission
- 5. Brain Atrophy

Diagnosis

- 1. Neurological Examination
- 2. Genetic Testing
- 3. Neuropsychological Assessment
- 4. Imaging Studies
- 5. Biomarker Testing





Table 1: Biomarkers of Neurodegenerative Diseases

| Biomarker name | Alzheimer's Disease (AD) | Parkinson's Disease (PD) | Huntington's Disease (HD) |
|---|--------------------------------|--------------------------------|---------------------------------|
| Phospho-tau181 (p-tau181) | √ | | |
| Phospho-tau217 (p-tau217) | √ | | |
| Phospho-tau231(p-tau231) | √ | | |
| Phospho-tau212 and 214 (p-tau212 and 214) | ~ | | |
| Phospho-tau202 and 205 (p-tau202 and 205) | √ | | |
| Phospho-tau413 (p-tau413) | √ | | |
| Phospho-tau422 (p-tau422) | √ | | |
| Tau proteins (Tau) | √ | √ | |
| Neurofilament light chain (NfL) | √ | √ | √ |
| Glial fibrillary acid protein (GFAP) | √ | √ | |
| Beta-amyloid 42 (Aβ42) | √ | √ | |
| Beta-amyloid 38 (Aβ38) | √ | | |
| Beta-amyloid 40 (Aβ40) | √ | √ | |
| Monocyte chemoattractant protein-1 (MCP-1/CCL2) | √ | | √ |
| Neurogranin protein (Neurogranin) | √ | | |
| Neuron-specific enolase (NSE) | √ | | |
| sTREM2 | √ | | |
| Visinin-like protein 1 (VLP-1) | ~ | | |
| Chitinase-3-like protein 1 (CHI3L1/YKL-40) | ~ | | |
| Alpha-synuclein (α-synuclein) | √ | √ | |



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| Apolipoprotein E (APOE4) | V | | |
|---|--------------|---|--------------|
| AD-associated thread protein (AD7c-NTP) | \checkmark | | |
| Human Parkinsonism associated deglycase (PARK7) | | V | \checkmark |
| Brain-derived neurotrophic factor (BDNF) | | | \checkmark |
| Interleukin-6 (IL-6) | | | |
| Pro-neuropeptide Y | | | \checkmark |
| Heart fatty acid-binding protein (HFABP) | V | | |

Biomarkers for Alzheimer's Disease

Table 2: Key biomarkers in blood and their detection limits [10-16].

| | Biomarker | Limit |
|---------------------------|---|---------------------------------------|
| | | (Pathological concentration in blood) |
| | Phospho-tau181 (p-tau181) | >3 pg/ml |
| | Phospho-tau217 (p-tau217) | >10 pg/ml |
| | Phospho-tau231(p-tau231) | >10 pg/ml |
| | Phospho-tau212 and 214 (p-tau212 and 214) | 1 |
| | Phospho-tau202 and 205 (p-tau202 and 205) | 1 |
| | Phospho-tau413 (p-tau413) | 1 |
| | Phospho-tau422 (p-tau422) | 1 |
| Alzheimer Disease (AD) | Tau proteins (Tau) | >100 pg/ml |
| | Neurofilament light chain (NfL) | >30 pg/ml |
| | Glial fibrillary acid protein (GFAP) | >100 pg/ml |
| | Beta-amyloid 42 (Aβ42) | 1 |
| | Beta-amyloid 38 (Aβ38) | 1 |
| | Beta-amyloid 40 (Aβ40) | 1 |
| | Monocyte chemoattractant protein-1 (MCP-1/CCL2) | 1 |
| | Neurogranin protein (Neurogranin) | 1 |
| | Neuron-specific enolase (NSE) | 1 |
| | sTREM2 | 1 |
| | Visinin-like protein 1 (VLP-1) | 1 |



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| Chitinase-3-like protein 1 (CHI3L1/YKL-40) | 1 |
|---|---|
| Alpha-synuclein (α-synuclein) | 1 |
| Apolipoprotein E (APOE4) | 1 |
| AD-associated neuaronal thread protein (AD7c-NTP) | 1 |
| Heart fatty acid-binding protein (HFABP) | 1 |

Sample Types: Blood, cerebrospinal fluid (CSF)

Biomarkers for Parkinson's Disease

Table 3: Key biomarkers in blood and their detection limits.

| | Biomarker | Limit (Pathological concentration in blood) |
|------------------------------|---|---|
| | Human Parkinsonism associated deglycase (PARK7) | 1 |
| | Alpha-synuclein (α-synuclein) | >10 pg/ml |
| Parkinsonism Disease (PD) | Tau proteins (Tau) | >10 pg/ml |
| | Neurofilament light chain (NfL) | I |
| | Glial fibrillary acid protein (GFAP) | 1 |
| | Beta-amyloid 42 (Aβ42) | 1 |
| | Beta-amyloid 40 (Aβ40) | >100 pg/ml |

Sample Types: Blood and cerebrospinal fluid (CSF)



Biomarkers for Huntington's Disease

| Table 4: Key | y biomarkers | in | blood ar | nd their | detection | limits. |
|--------------|--------------|----|----------|----------|-----------|---------|
| | | | | | | |

| Biomarker | | Limit (Pathological concentration in blood) |
|---|--|--|
| | Brain-derived neurotrophic factor (BDNF) | I |
| Huntington's Disease (HD) Biomarker | Monocyte chemoattractant protein-1 (MCP-1/CCL2) | J |
| | Interleukin-6 (IL-6) | I |
| | Neurofilament light chain (NfL) | 1 |
| | Pro-neuropeptide Y | 1 |

Sample Types: Blood, cerebrospinal fluid (CSF)

3. Technologies and Platforms for Neurodegenerative Diseases Diagnostics

ELISA (Enzyme-Linked Immunosorbent Assay)

- Overview: It is a commonly used detection technology that determines the presence or absence of specific proteins or other molecules contained in a sample and combines antibodies and an enzyme label.
- Advantages: High sensitivity, strong specificity, simple operation, and relatively low cost.

Simoa (Single Molecule Array)

- Overview: Unlike traditional ELISA techniques, this technology can detect very low levels of proteins, nucleic acids and other biomolecules at a sensitivity several orders of magnitudes greater. Through using an array of sealed microwells and fluorescently labeled single molecules, Simoa is able to achieve detection and quantification of individual molecules. This capability is essential for early disease diagnosis, biomarker detection, scientific research ^[17-19].
- Advantages:
 - (1) Ultra-high Sensitivity: Simoa can detect biomarkers at extremely low concentrations, even at the



single-molecule level, which is several orders of magnitude more sensitive than traditional methods such as ELISA.

- (2) Early Disease Detection: Due to its high sensitivity, Simoa technology can detect trace biomarkers at very early stages of diseases, facilitating early diagnosis and treatment.
- (3) Accurate Quantification: Simoa not only detects the presence of biomarkers but also accurately quantifies their concentrations, which is crucial for precise disease monitoring and treatment.
- (4) Versatility and Flexibility: The application of this technology extends to other biomolecules, such as proteins and nucleic acids, thus it is extremely valuable for a wide range of biomedical research and clinical applications.
- (5) Reduced Sample Volume: Simoa requires only a very small amount of sample for testing, reducing the need for large sample volumes from patients.
- (6) High Throughput and Automation: The Simoa platform supports high-throughput sample analysis and can be automated, improving the efficiency and reproducibility of experiments.



Figure: Detecting misfolded proteins in blood through SIMOA technology for the diagnosis of Alzheimer's disease / Source: researchgate.net

Multiplex Assays

• **Overview:** Simultaneous detection of multiple biomarkers with enhanced detection efficiency and increased data throughput achieved with different fluorescent or chemically labels specifying each biomarker.

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• Advantages: High throughput, sample and reagent savings, strong data integration.

ECL (Electrochemiluminescence)

- **Overview:** A high-sensitivity detection method that combines electrochemical and luminescence technologies to analyze biomarkers and molecular interactions.
- Advantages: Extremely high sensitivity and specificity, low background noise, suitable for detecting low-concentration samples and high-throughput screening.

LC-MS/MS (Liquid Chromatography-Tandem Mass Spectrometry)

- **Overview:** A high-precision detection method that combines liquid chromatography and mass spectrometry to analyze compounds in complex biological samples.
- **Advantages:** High sensitivity and specificity, capable of detecting multiple target molecules simultaneously, suitable for metabolomics and proteomics research.

ECLIA (Electrochemiluminescence Immunoassay)

- Overview: A combined electrochemical-luminescence immunoassay method for biomarker analysis.
- Advantages: Extremely high sensitivity and specificity, suitable for detecting low-concentration samples, low background noise.

CLEIA (Chemiluminescent Enzyme Immunoassay)

- **Overview:** A chemiluminescence enzyme linked immunoassay (CLIEA) method for detection of biomarkers through a chemical reaction that in a light signal.
- Advantages: High sensitivity, high specificity, rapid detection, suitable for analyzing various sample types.

4. Clinical Applications and Research in Alzheimer's Disease Diagnostics

| Name | Sample | Biomarkers | Method | |
|------------------------------|--------|----------------------------------|----------|--|
| | Blood | P-tau217 | | |
| Roche | CSF | P-tau181/Aβ42 ratio | ECL | |
| Blood | | P-tau181+APOE E4 | | |
| C ₂ N Diagnostics | Blood | Aβ42/40 ratio and p-tau217 ratio | LC-MS | |
| | | Αβ42/40 | LC-MS/MS | |
| Quest Diagnostics | Blood | Blood p-tau217 | | |
| | | p-tau181 | IA | |



| | | p-tau181 | |
|-----------|----------|----------|-------|
| Quanterix | Blood | NfL | Simoa |
| | | p-tau217 | |
| Alzpath | Blood | p-tau217 | Simoa |
| | p-tau217 | | |
| | | Αβ42/40 | |
| Labcorp | Blood | GFAP | CLEIA |
| | | p-tau181 | |
| | | NfL | |

5. GeneMedi's Alzheimer's Disease Diagnostics Solution

| | Name | Cat No. |
|---|---|--------------------|
| | Phospho-tau181 (p-tau181) | GMP-h-p-tau181 |
| | Phospho-tau217 (p-tau217) | GMP-h-p-tau217 |
| | Phospho-tau231(p-tau231) | GMP-h-p-tau231 |
| | Phospho-tau212 and 214 (p-tau212 and 214) | GMP-h-p-tau212/214 |
| | Phospho-tau202 and 205 (p-tau202 and 205) | GMP-h-p-tau202/205 |
| | Phospho-tau413 (p-tau413) | GMP-h-p-tau413 |
| | Phospho-tau422 (p-tau422) | GMP-h-p-tau422 |
| | Tau proteins (Tau) | GMP-h-Tau |
| | Neurofilament light chain (NfL) | GMP-h-NfL |
| Genemedi's Alzheimer's disease-related biomarker | Glial fibrillary acid protein (GFAP) | GMP-h-GFAP |
| | Beta-amyloid 42 (Aβ42) | GMP-h-Aβ42 |
| | Beta-amyloid 38 (Aβ38) | GMP-h-Aβ38 |
| products | Beta-amyloid 40 (Aβ40) | GMP-h-Aβ40 |
| | Monocyte chemoattractant protein-1 (MCP-1/CCL2) | GMP-h-MCP-1 |
| | Neurogranin protein (Neurogranin) | GMP-h-Neurogranin |
| | Neuron-specific enolase (NSE) | GMP-h-NSE |
| | sTREM2 | GMP-h-sTREM2 |
| - | Visinin-like protein 1 (VLP-1) | GMP-h-VLP-1 |
| | Chitinase-3-like protein 1 (CHI3L1/YKL-40) | GMP-h-YKL-40 |
| | Alpha-synuclein (α-synuclein) | GMP-h-α-synuclein |
| | Apolipoprotein E (APOE4) | GMP-h-APOE4 |
| | AD-associated neuaronal thread protein (AD7c-NTP) | GMP-h-AD7c-NTP |
| | Heart fatty acid-binding protein (HFABP) | GMP-h-HFABP |

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 GeneMedi's three phosphorylated tau protein antibodies (p-tau181, p-tau217, and p-tau231) demonstrate outstanding sensitivity and exceptional specificity with their corresponding phosphorylated forms in direct ELISA assays.



- Fig 1. Validation of GeneMedi's Anti-human p-tau181/p-tau217/p-tau231 antibody.
- GeneMedi's three phosphorylated tau protein antibodies (p-tau181, p-tau217, and p-tau231) show excellent sensitivity and extreme specificity with their corresponding phosphorylated forms in direct ELISA assays.



| Antibody | GMP-h-p-tau217-Ab01 | | | | |
|----------|----------------------------------|----------------------------------|----------------------------------|---------------------------------------|---------------------------------|
| Antigen | GMP-h-p-tau181- Ag01(peptide) | GMP-h-p-tau217- Ag01(peptide) | GMP-h-p-tau231- Ag01(peptide) | GMP-h-p-tau217- Ag02 (full length) | GMP-h-Tau- Ag01(full length) |
| EC50 | 27320 ng/ml | 4.326 ng/ml | 3563 ng/ml | 6.943 ng/ml | 1843 ng/ml |

Fig 2. Direct ELISA Binding Curves of Phosphorylated Tau217 Antibody with Various Antigens.

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3) GeneMedi's three phosphorylated tau protein antibodies (p-tau181, p-tau217, and p-tau231) display outstanding sensitivity and exceptional specificity to their corresponding phosphorylated forms in direct ELISA assays.



| Antigen | GMP-h-p-tau217-Ag02 (full length) | | |
|---|-----------------------------------|-----------------------------|--|
| Antibody pair Coating-GMP-h-p-tau217-Ab01 | | Coating-GMP-h-p-tau217-Ab01 | |
| EC50 | 76.55 ng/ml | 39.02 ng/ml | |





| Antigen | GMP-h-tau-Ag01 (full length) | | |
|---------------|------------------------------|-----------------------------|--|
| Antibody pair | Coating-GMP-h-p-tau217-Ab01 | Coating-GMP-h-p-tau217-Ab01 | |
| | Detection-GMP-h-Tau-Ab01 | Detection-GMP-h-Tau-Ab03 | |
| EC50 | 4352000 ng/ml | 000 ng/ml 20030 ng/ml | |



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| Antigen | GMP-h-p-tau217-Ag01 (peptide) | | | | |
|---------------|---------------------------------|---------------------------------|---------------------------------|---------------------------------|--|
| Antibody pair | Coating-GMP-h-p-tau 217-Ab01 | Coating-GMP-h-p-tau 217-Ab01 | Coating-GMP-h-p-tau 217-Ab02 | Coating-GMP-h-p-tau 217-Ab02 | |
| | Detection-GMP-h-Tau -Ab01 | Detection-GMP-h-Tau -Ab02 | Detection-GMP-h-Tau -Ab01 | Detection-GMP-h-Tau -Ab02 | |
| EC50 | NA | NA | NA | NA | |

Fig 3. Sandwich ELISA Analysis of Full-Length Phosphorylated Tau 217 Antigen with Corresponding Phosphorylated Tau 217 Antibody and Total Tau Antibody.



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PRODUCT LIST

| Cat.No | Products Name | Products Type |
|--------------------|---|---------------|
| GMP-h-p-tau181 | Human Phospho-tau181 (p-tau181) antigen/antibody | peptide |
| GMP-h-p-tau217 | Human Phospho-tau217 (p-tau217) antigen/antibody | peptide |
| GMP-h-p-tau231 | Human Phospho-tau231 (p-tau231) antigen/antibody | peptide |
| GMP-h-p-tau212/214 | Human Phospho-tau212 and 214 (p-tau212 and 214) antigen/antibody | peptide |
| GMP-h-p-tau202/205 | Human Phospho-tau202 and 205 (p-tau202 and 205) antigen/antibody | peptide |
| GMP-h-p-tau413 | Human Phospho-tau413 (p-tau413) antigen/antibody | peptide |
| GMP-h-p-tau422 | Human Phospho-tau422 (p-tau422) antigen/antibody | peptide |
| GMP-h-Tau | Human tau proteins (Tau) antigen/antibody | protein |
| GMP-h-NFL | Human neurofilament light chain (NFL) antigen/antibody | protein |
| GMP-h-GFAP | Human Glial fibrillary acid protein (GFAP) antigen/antibody | protein |
| GMP-h-Aβ42 | Human beta-amyloid 42 (Aβ42) antigen/antibody | peptide |
| GMP-h-Aβ38 | Human beta-amyloid 38 (Aβ38) antigen/antibody | peptide |
| GMP-h-Aβ40 | Human beta-amyloid 40 (Aβ40) antigen/antibody | peptide |
| GMP-h-MCP-1 | Human Monocyte chemoattractant protein-1 (MCP-1/CCL2) antigen/antibody | peptide |
| GMP-h-neurogranin | Human neurogranin protein (neurogranin) antigen/antibody | protein |
| GMP-h-NSE | Human neuron-specific enolase (NSE) antigen/antibody | protein |
| GMP-h-sTREM2 | Human soluble fragment of triggering receptor expressed on myeloid cells 2 (sTREM2) antigen/antibody | protein |
| GMP-h-VLP-1 | Human Visinin-like protein 1 (VLP-1) antigen/antibody | protein |
| GMP-h-YKL-40 | Human Chitinase-3-like protein 1 (CHI3L1/YKL-40) antigen/antibody | protein |
| GMP-h-a-synuclein | Human Alpha-synuclein (α-synuclein) antigen/antibody | protein |
| GMP-h-APOE4 | Human Apolipoprotein E (APOE4) antigen/antibody | protein |
| GMP-h-AD7c-NTP | Human AD-associated neuaronal thread protein (AD7c-NTP) antigen/antibody | protein |



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PRODUCT LIST

| Cat.No | Products Name | Products Type |
|----------------------|--|---------------|
| GMP-h-PARK7 | Human Parkinsonism associated deglycase (PARK7) antigen/antibody | protein |
| GMP-h-BDNF | Human brain derived neurotrophic factor (BDNF) antigen/antibody | protein |
| GMP-h-Neuropeptide-Y | Human Pro-neuropeptide Y (Neuropeptide Y) antigen/antibody | protein |
| GMP-h-HFABP | Human Heart fatty acid-binding protein (HFABP) antigen/antibody | protein |

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GeneMedi specializes in creating superior antibody, protein, and vectorbased bioproducts, revolutionizing diagnostics and therapeutic solutions.

At GeneMedi, innovation, product integrity, and scalable solutions form the cornerstone of our mission to advance the field of diagnostics and therapeutics. Our portfolio of antibodies, proteins, and vector-derived products is built on a foundation of unparalleled expertise in the following areas:

Innovative Antigen Design and Robust Assay Development

Our strategic focus on biomarkers and targets analysis enables the creation of highly specific antigens and the development of robust assays, ensuring our products achieve superior performance in clinical and research settings.

Streamlined Molecular Discovery with Emphasis on Stability

Rapid Protein & Antibody Identification: Our proprietary platforms, TAURUS for accelerated antibody discovery and LIBRA for Al-driven protein evolution, are designed to identify and optimize molecules with optimal stability and functionality.

Cutting-Edge AAV & GCT Discoveries: The G-NEXT platform is our answer to the industry's need for innovative AAV vectors, offering improved stability, efficiency, and safety forgroundbreaking gene therapy approaches.

Scalable Production and Uncompromising Quality

High-Volume Protein & Antibody Manufacturing: Our facilities are equipped to handle large-scale production up to 1000L per batch, ensuring high levels of purity and stability through stringent quality controls.

Advanced Vector Manufacturing Capabilities: With a focus on AAV, Lentivirus, and VLP production up to 200L per batch, we employ sophisticated purification techniques to guarantee vector efficacy and integrity

Comprehensive Solutions for Diverse Application Needs

Diagnostics: Our diagnostic solutions leverage CLIA, LFA, and ELISA platforms for comprehensive assay validation and clinical sample consistency, setting new standards in diagnostic accuracy.

Therapeutics: We specialize in analytical development for ADCs, AAV therapies, and CAR-X technologies, ensuring our therapeutic products are characterized by their specificity, potency, and safety.

GeneMedi (GM) is dedicated to delivering innovative and scalable biotechnological solutions, driving forward the fields of diagnostics and therapeutics with confidence and expertise. Contact with GM to reach your reliable industrial partner.

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