

The Evolution of Gene Therapy Drug Development

$$2 + 2 = 5$$

Why?

Innovative Synergy for a
Supra-additive Effect!

Founded in 2015 in London (UK), Cell and Gene Therapy Ltd. has emerged as a leading biotech company and a principal member of an international group specializing in the development of gene therapy drugs. The core concept of the products lies in the masterful combination of traditional and innovative elements, creating a supra-additive effect and forging a unique solution. Within this concept, the following components are envisioned:

Unique DNA Vectors

Our DNA vectors, developed and patented as a platform solution, elegantly combine efficacy, safety, and flexibility by integrating universal or tissue-specific and inflammation-activated promoters with coding sequences of target genes.

Multiplicity of Targets

Therapeutic diversity is achieved through the use of a composition of next-generation DNA vectors containing genes, each of which is directed at corresponding target.

Advanced Delivery Systems

The use of modern delivery systems enables the attainment of therapeutically significant concentrations of target proteins, thereby maximizing the drug's therapeutic effect.

Technological Excellence

The use of proven technologies allows the drug to be manufactured at various standard biotechnological facilities, achieving both competitive pricing and high profitability.



Our development priorities focus on diseases that currently have no effective treatments available, such as **Alzheimer's disease, Parkinson's disease, multiple sclerosis, liver fibrosis, myocardial fibrosis**, along with numerous other diseases. Furthermore, we are dedicated to addressing **type 2 diabetes mellitus and obesity**, as well as rare and orphan diseases.

Focus on Pathology

Priority is given to the careful selection of targets within pathological biological processes, focusing on underlying mechanisms rather than merely addressing disease symptoms, to achieve effective and sustainable therapeutic outcomes.

Use of Native Genes

The use of native genes ensures harmonious integration with natural biological processes, reducing the risk of adverse reactions and enhancing the drug's biocompatibility.

Precision Delivery

The use of optimal promoters ensures accurate and efficient delivery of DNA vectors to target cells, enhancing overall therapy efficacy and minimizing off-target effects.

Regulatory Compliance

The vectors' structural elements, developed as part of our platform solution, fully comply with FDA and EMA requirements, ensuring strict safety and efficacy standards.

Since 2015, an international team of scientists, spearheaded by our company, has dedicated extensive intellectual resources and cutting-edge research efforts to this project, ultimately leading to the development of a groundbreaking universal platform solution — non-viral DNA vectors series VTvaf17 and GDDT1.8NAS — for creating advanced genetic tools in the rapidly evolving fields of biomedical and genetic technologies. These DNA vectors incorporate the unique RNA-out regulatory element from the Tn10 transposon, thus enabling antibiotic-free positive selection, and offering the following key advantages:

Maximum Safety

The **absence of antibiotic resistance genes** and **viral genome sequences** in our DNA vectors, in accordance with EMA and FDA recommendations, ensures the highest safety. This distinct combination in a non-viral DNA vectors makes our solution **one-of-a-kind globally**.

Nature-like Mechanism

The use of **non-modified native genes** ensures seamless integration with natural biological processes, minimizing the risk of adverse reactions.

Precision Expression

By integrating **cell-specific and inflammation-activated promoters**, our drugs achieve precise and effective expression of genes in target cells while minimizing undesirable side effects.



**DNA vectors
VTvaf17,
GDDT1.8NAS**



Superior Performance

By incorporating **advanced delivery systems** into our drug, we achieve therapeutically significant concentrations of target proteins.

Multiply Therapeutic Targets

Creation of a **unique composition of genes** empowers our drugs to simultaneously target multiple therapeutic pathways, achieving a synergistic effect.

Technological Excellence

Implementation of high-tech manufacturing techniques optimizes production processes, achieving **exceptional efficiency** and **significant cost reductions**.



The intellectual property associated with this project is protected by **more than 30 patents** across various countries worldwide, highlighting the **unique** and **innovative** nature of the product.

We have developed the unique Alzheimer's gene therapy candidate **CG-AD211** that clears toxic tau aggregates, stimulates neuroregeneration, and protect neurons—all tasks that require orchestrating many genes, yet have traditionally carried risks for healthy cells. This is a challenge for conventional pharmacology but we overcome it by using a) composite promoter, which activates only in the target cells and only under inflammation, b) two-stage administration regimen, c) a meticulously selected gene set, thereby delivering therapy exactly where it is needed while eliminating risks to healthy cells.

Focus on Pathology

Priority attention is given to the disposal of pathological intracellular protein aggregates and to the processes of neuroregeneration of damaged neurons to restore bodily functions.

Multiplicity of Targets

Therapeutic versatility is achieved through the use of a composition of genes responsible for a range of biological processes — including the disposal of pathological protein aggregates, neuron restoration, and the regeneration of synaptic connections.

Advanced Delivery Systems

A complex of cationic liposomes + PEG with *** functionalization's been selected as the delivery system, which ensures the achievement of therapeutically significant concentrations of target proteins, thereby maximizing the therapeutic effect.

Technological Excellence

The use of proven technologies allows the drug to be manufactured at various standard biotechnological facilities, achieving both competitive pricing and profitability.

Treatment Protocols

Two-stage administration: The drug is administered sequentially, with each stage targeting a specific goal: 1. Disposal of pathological intracellular protein aggregates; 2. Neuroregeneration; which ensures maximum drug efficacy and minimizes adverse effects.

Unique DNA Vectors

Our therapeutic DNA vectors of GDDT1.8NAS series developed and patented as a platform solution, elegantly combine efficacy, safety, and the flexibility to vary with universal and tissue-specific promoters and coding sequences of genes.

Use of Native Genes

The use of native genes H***, H***, C***, L***, S***, L***, T***, A***, C***, B**, D**, R**, N**, C** ensures harmonious integration with natural biological processes, thereby reducing the risk of adverse reactions and enhancing the drug's biocompatibility.

Precision Expression

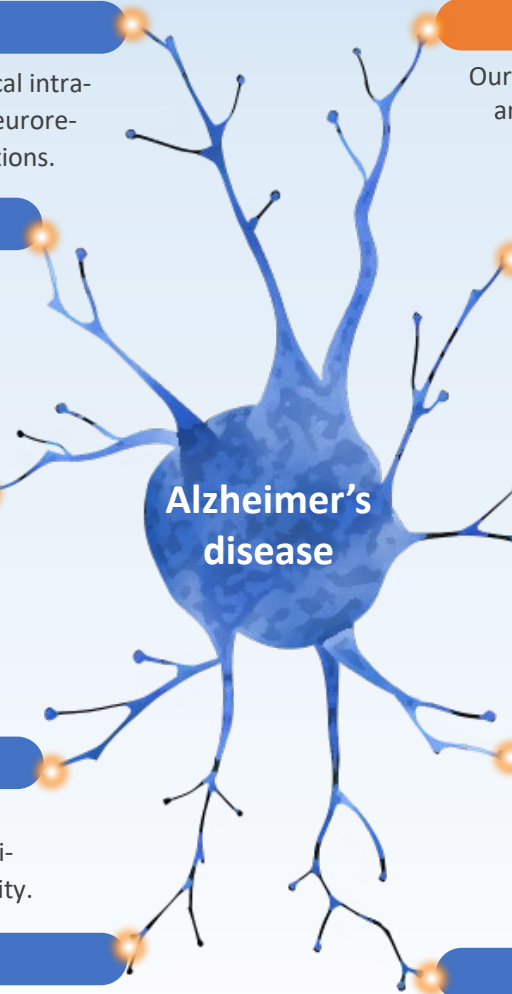
The use of combinations of tissue-specific and inflammation-activated promoters ensures precise and effective expression of delivered genes in target cells, that enhancing overall therapeutic efficacy and consequently minimizing ectopic effects.

Regulatory Compliance

The composition of structural elements of the vectors, developed and patented as part of a platform solution, fully complies with FDA and EMA requirements, guaranteeing adherence to strict safety and efficacy standards.

Method of administration

The intraarterial infusion route, combined with the preventive administration of mannitol, intravenous hydration, and ultrasound (FUS and UTMD), is aimed at ensuring maximum penetration of the delivered drug into the target tissues of the patient's brain.



The project's strategy involves the creation of a pharmaceutical agent for the therapeutic implementation of the following biological processes directly related to the disease, for which there are currently no registered medications. A treatment regimen - two-stage pulse administration - was selected to maximize drug efficacy while minimizing uncontrolled therapeutic-gene interactions:

Therapeutic Stage 1

Disposal of pathological intra-cellular protein aggregates

This stage is primarily focused on the elimination of tau aggregates. The active degradation system will ensure the clearance of other pathological protein aggregates (PPA), including amyloid beta conglomerates, TDP-43 protein, intracellular polyamides, etc.

Task	Description
Ubiquitination	Placing a tag for the recognition and targeting of PPA
Binding and transport	Adapter proteins recognize and bind PPA, directing them to autophagosomes for degradation
Autophagosome formation	Enhancing autophagy through autophagosome formation promotes effective degradation
Autophagolysosome formation	Fusion of the autophagosome with a lysosome ensures the final degradation of PPA and damaged organelles
Degradation and exocytosis	This contributes to maintaining cellular homeostasis and prevents neuronal damage

Therapeutic Stage 2

Regeneration of damaged regions of the brain

Neurons that have been compromised by the progressive accumulation of pathological intracellular protein aggregates indisputably require not merely protection but concerted regenerative intervention. Consequently, the brain's normal function will be reestablished.

Task	Description
Stabilization of microtubules	Stable microtubules allow neurons to properly reintegrate into the existing neural networks and restore synaptic connections
Axonal navigation	Directed growth of axons and dendrites to specific target cells is necessary for the restoration of synaptic connections
Synaptic connections	Neurons must re-establish functional synapses with other neurons to ensure effective transmission of nerve impulses
Neuroprotection	Slows the subsequent progression of neurodegenerative changes in the brain and prevents neuronal apoptosis

Exposure Period