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# Novel CRISPR Gene Therapy for Haploinsufficiency

Tech ID: 30127 / UC Case 2017-040-0

# **INVENTION NOVELTY**

This technology presents a way to treat human genetic disease caused by haploinsufficiency and reduced protein production. The method employs the use of adeno-associated viral (AAV) vectors for the in vivo delivery of a CRISPR-based gene expression activator (CRISPRa) that boosts transcription from the existing functional copy of the affected gene.

# **VALUE PROPOSITION**

Haploinsufficiency – having only one functional copy of a gene due to gene deletion or other gene-inactivating mutations, is the basis of many human genetic disorders caused by the insufficient amounts of product generated by the affected gene. Attempts to treat such developmentally-predetermined conditions have relied on gene therapy and the re-introduction of a second functioning gene copy. However, this therapeutic approach has been hampered by a series of confounding factors including gene size, mutations caused by gene or vector insertion, and inability to target specific cells or deliver specific doses of the gene. A unique merit of this novel technology is the use of non-pathogenic and non-integrating AAV vector to deliver a CRISPRa, which is targeted to the enhancer or promoter region of a gene. In the case of low gene dosage-based conditions, this allows for inducing transcription from the existing copy of the gene without creating any changes in the patient's DNA. Furthermore, as the AAV vector will not be carrying the actual missing gene copy, this method circumvents common limitations to using AAV for effective delivery in patients.

## **TECHNOLOGY DESCRIPTION**

Gene transcription is increased by a nuclease defective Cas9 (dCas9) fused to a transcriptional activation domain combined with a unique guide RNA construct targeted to the regulatory (promoter and/or enhancer) regions of a gene. Thus, this CRISPRa complex does not cause nicks or breaks in the DNA, can be easily packaged in an AAV vector for in vivo delivery, and cell/tissuespecificity is determined by the targeted cis-regulatory regions. The inventors have demonstrated successful in vivo use of the technology by curing obesity in Sim1 and Mc4r mouse haploinsufficiency models, which as is the case in humans, results in defects in the leptin pathway and hyperphagic obesity.

## LOOKING FOR PARTNERS

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# **INVENTORS**

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**OTHER INFORMATION** 

**KEYWORDS** Haploinsufficiency, Gene

therapy, CRISPR

#### **CATEGORIZED AS**

Medical Disease: Genetic **Diseases and Dysmorphic Syndromes** Gene Therapy **RELATED CASES** 

2017-040-0

To develop & commercialize the technology as a gene therapy tool for the treatment of human disease based on genetic

haploinsufficiency.

# **STAGE OF DEVELOPMENT**

Pre-clinical / Proof of concept

#### **RELATED MATERIALS**

Matharu, N., Rattanasopha, S., Tamura, S., Maliskova, L., Wang, Y., Bernard, A., ... & Ahituv, N. (2019).
CRISPR-mediated activation of a promoter or enhancer rescues obesity caused by haploinsufficiency.
Science, 363(6424), eaau0629.

## **OTHER INFORMATION**

Commentary: Montefiori, L. E., & Nobrega, M. A. (2019). Gene therapy for pathologic gene expression. Science, 363(6424), 231-

#### 232.

# **DATA AVAILABILITY**

Under CDA / NDA

#### **PATENT STATUS**

Country	Туре	Number	Dated	Case
China	Issued Patent	110612113	03/26/2024	2017-040
United States Of America	Issued Patent	11,730,828	08/22/2023	2017-040
European Patent Office	Published Application	3579858	12/18/2019	2017-040
Australia	Published Application 2017-040			
Japan	Published Application 2017-040		2017-040	

Additional Patents Pending

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