

Delivering on the promise of Prime Editing

LNP delivered Prime Editors restore glycemic control in humanized rodent models of Glycogen Storage Disease Type 1b (GSD1b)

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On behalf of the team at Prime Medicine

Disclosures

Vivian Choi declares she is currently an employee of Prime Medicine, Inc. and owns equity in Prime Medicine.



Prime Editing is programmable for both search and replace

The PE technology utilizes a Prime Editor protein and a Prime Editing guide RNA (pegRNA) to directly write new genetic information into a targeted DNA site without requiring a DSB



We believe Prime Editing is the only gene editing technology that medicine can edit, correct, insert and delete DNA sequences in any target tissue

Corrects mutations across many tissues, organs and cell types, in dividing and non-dividing human cells



Prime Editing platform modularity accelerates and de-risks ongoing efforts, enabling rapid generation of new product candidates

Core components can be readily leveraged to accelerate pipeline growth, efficiency and execution



Prime Medicine's approach to developing Prime Editors to treat liver and metabolic diseases

Prime Editors are specific to *human* patient DNA sequence and designed for the correction of *human* mutations



- Establish potency of lead Prime Editor drug candidates
- Establish genotype to phenotype correlation and off-target profile
- Establish pharmacology, safety, tolerability
- Determine biodistribution, drug pharmacokinetics
- Determine PK/PD relationships, human dose projections

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Prime has developed a universal LNP for our liver & metabolic programs

Α

Prime Medicine's Universal LNP contains a novel GalNAc targeting ligand

Shared LNP/PE components



✓ Delivery to the liver via the ASGPR is a validated delivery mechanism

Compared to LNPs without a targeting ligand, Prime's Universal LNP*:

- ✓ Increases potency
- ✓ Improves safety profile
- ✓ Improves biodistribution

Prime Medicine's modular LNP can be used to generate multiple different drug product candidates (DP)





By swapping only the guide RNAs while keeping the other components constant, we have a new product with the potential for the same critical quality attributes

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Therapeutic approach: LNP-mediated delivery of Prime Editor components to liver

One-time delivery of LNP Prime Editor cargo with potential to correct pathogenic mutations in the liver





Prime Editors to correct pathogenic mutations causing von Gierke disease or Glycogen Storage Disease Type 1b (GSD1b)

Initially correct the two most prevalent mutations that cause GSD1b, carried by ~ 50% of patients

Glycogen Storage Disease Type 1b (GSD1b)

Description:

• Rare disease preventing production of glucose during fasting state, causing hypoglycemic episodes and associated seizures

Human genetics and biology:

- Autosomal recessive, caused by mutations in the *SLC37A4* gene that encodes G6PT, a glucose-6-phosphate transporter
- SLC37A4 p.L348fs and p.G339C mutations found in ~50% of GSD1b patient population

Unmet need:

- Standard of care focuses on reducing symptom severity rather than treating the underlying cause of the disease
- No disease modifying therapies approved

Prime Medicine's approach:

 IV administration of liver targeted LNP Prime Editors to correct either the p.L348fs or p.G339C mutations to restore glucose homeostasis in patients with GSD1b



Identification of lead Prime Editors for correction of the p.G339C and p.L348fs mutations



Prime Medicine's high throughput screening platform identifies multiple Prime Editors capable of efficient correction of p.G339C and p.L348fs in primary mouse hepatocytes isolated from GSD1b humanized mice

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Optimization of components improves precise correction of the p.L348fs mutation *in vivo*



Improved in vivo correction observed following optimization of Prime Editors components



Unintended editing rates are below 0.2% at every dose tested

*Based on PK/PD relationships and quantification of cell types in liver: Wang et al Sci. Rep. (2021) 11:19396; MacParland et al Nat Commun. (2018) 9:4383; Hansel et al, Curr Protoc Toxicol (2014) 62:14.12.1; 1minec, Adv Anat Embryol Cell Biol. (2001) 161:III-XIII. 1-151. ** Calculation based on 60% of cells in whole liver are hepatocytes; # Unintended edits = any SNVs or indels within 300bp either side of the edit site.

Optimization of components improves precise correction of the p.G339C mutation *in vivo*



Improved in vivo correction observed following optimization of Prime Editors components



*Based on PK/PD relationships and quantification of cell types in liver: Wang et al Sci. Rep. (2021) 11:19396; MacParland et al Nat Commun. (2018) 9:4383; Hansel et al, Curr Protoc Toxicol (2014) 62:14.12.1; Kmiec, Adv Anat Embryol Cell Biol. (2001) 161:III–XIII. 1–151. ** Calculation based on 60% of cells in whole liver are hepatocytes; # Unintended edits = any SNVs or indels within 300bp either side of the edit site.



G6PT transcripts and protein are restored in humanized p.L348fs mouse liver following delivery of Prime Editors

Precise editing of the G6PT gene restores G6PT protein expression



Prime Editing restores glucagon-dependent glucose production in a mouse model of GSD1b







Prime editing improves metabolic dysfunction observed in a mouse model of GSD1b





Successful Prime Editing with precise correction leads to **reduction of fasting hypoglycemia, triglycerides, liver glycogen & restores glucagon responsiveness** in humanized mice at a dose predicted to be clinically relevant for human disease

Prime Editing in NHP



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Prime Medicine's Universal LNP exhibits an excellent safety profile in cynomolgus monkey (NHP)

Well-tolerated with no acute reactions, clinical observations, or body weight changes
Animals healthy at 54 weeks



- No observed change in platelets, coagulation time or blood count
- No observed change in blood biochemistry panel



- Minimal changes in serum IL6 levels
- No other observed cytokine changes

No changes observed in liver histopathology (H&E) Minimal transient LFT elevations

Benchmarked against other LNPs in clinical development

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Safety



Preliminary analysis: no detectable off-target editing in patient cells treated with Prime Editor



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¹Targeted Analysis of potential off-target sites using targeted deep sequencing in Prime Edited human patient iPSC cells. GSD1b: glycogen storage disease type 1b, ATP7B: ATPase copper transporting beta, G6PT: glucose-6-phosphate translocase, Indels: insertions/deletions



Summary

Modular LNP platform

Prime has developed a universal liver targeted LNP-PE platform with the potential to deliver Prime Editors to precisely correct disease-causing mutations

- GalNAc targeting ligand improves dose potency, editing, and biodistribution compared to LNPs without targeting ligand
- > Excellent and differentiated safety profile in large animal & rodent studies

Glycogen Storage Disease type Ib

- LNP-RNA Prime Editor candidates achieve 80-90% precise hepatocyte correction of the *SLC37A4* (G6PT) gene mutations p.L348fs and p.G339C in humanized mice at clinically relevant doses
 - > IV delivery Prime Editor restores hepatic glycogen metabolism in a humanized mouse model of GSD1b
 - Large animal cynomolgus monkey studies demonstrate up to 83% precise hepatocyte editing of G6PT gene at p.L348 using a NHP surrogate pegRNA at a dose that was safe, well tolerated and durable

Off-target editing

> No off-target editing was detected in human cells derived from either Wilson's Disease or GSD1b patients

Prime Medicine's presentation on Prime Editors development for Wilson's Disease Monday Nov 18th at 8:45am (Location: 6C – Session Genetic & Metabolic Liver Diseases)





