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## **AIMers**

Correcting driver mutations for the treatment of Alpha-1 antitrypsin deficiency (AATD) and beyond

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3rd RNA Editing Summit: April 5-7, 2022



# Forward looking statements

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# Unlocking RNA editing with PRISM<sup>™</sup> to develop GalNAc-AIMers: A-to-I editing oligonucleotides

Optimize AIMer design for endogenous transcripts and GalNAc conjugation



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### ADAR enzymes

- First publication (1995) using oligonucleotide to edit RNA with endogenous ADAR<sup>1</sup>
- Catalyze conversion of A-to-I (G) in doublestranded RNA substrates
- ADAR1 is ubiquitously expressed across tissues, including liver and CNS – sizeable target space
- Cellular reservoir of ADAR capacity supports directed editing in addition to homeostatic function

### **Optimize AIMer design**

- Substrate learnings from biology and structures
- Applied to oligonucleotides
- Applied PRISM chemistry

### **PRISM-driven gains**



<sup>1</sup>Woolf et al., 1995 Proc Natl Assoc Sci 92:8298-8302; Monian et al., 2022 Nature Biotech published online Mar 7, 2022 doi: 10.1038.s41587-022-01225-1

# Proof-of-concept RNA editing in NHP liver is robust, durable, and specific



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AIMer tissue concentration (left) and editing activity (middle) in liver after 5 mg/kg SC injections on days 1-5; Right: Hepatocytes treated with 1 µM AIMer, 48 hrs later RNA collected, RNAseq conducted using strand-specific libraries to quantify editing; plotted circles represent sites with LOD>3. NHP: non-human primate; ACTB: Beta-actin; Monian et al., 2022 published online Mar 7, 2022; doi: 10.1038.s41587-022-01225-1

# Proof-of-concept preclinical RNA editing data published in *Nature Biotechnology* (March 2022)

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## Endogenous ADAR-mediated RNA editing in non-human primates using stereopure chemically modified oligonucleotides

Prashant Monian <sup>12</sup>, Chikdu Shivalila <sup>12</sup>, Genliang Lu<sup>1</sup>, Mamoru Shimizu<sup>1</sup>, David Boulay<sup>1</sup>, Karley Bussow<sup>1</sup>, Michael Byrne<sup>1</sup>, Adam Bezigian<sup>1</sup>, Arindom Chatterjee<sup>1</sup>, David Chew<sup>1</sup>, Jigar Desai<sup>1</sup>, Frank Favaloro<sup>1</sup>, Jack Godfrey<sup>1</sup>, Andrew Hoss<sup>1</sup>, Naoki Iwamoto<sup>1</sup>, Tomorni Kawamoto<sup>1</sup>, Jayakanthan Kumarasamy<sup>1</sup>, Anthony Lamattina<sup>1</sup>, Amber Lindsey<sup>1</sup>, Fangjun Liu<sup>1</sup>, Richard Looby<sup>1</sup>, Subramanian Marappan<sup>1</sup>, Jake Metterville<sup>1</sup>, Ronelle Murphy<sup>1</sup>, Jeff Rossi<sup>1</sup>, Tom Pu<sup>1</sup>, **Bijay Bhattarai**<sup>©</sup>, Stephany Standley<sup>1</sup>, Snehlata Tripathi<sup>1</sup>, Hailin Yang<sup>1</sup>, Yuan Yin<sup>1</sup>, Hui Yu<sup>1</sup>, **Cong Zhou**<sup>©</sup>, Luciano H. Apponi<sup>1</sup>, Pachamuthu Kandasamy<sup>1</sup> **and Chandra Vargeese<sup>®1</sup>** 

Technologies that recruit and direct the activity of endogenous RNA-editing enzymes to specific cellular RNAs have therapeutic potential, but translating them from cell culture into animal models has been challenging. Here we describe short, chemically modified oligonucleotides called AlMers that direct efficient and specific A-to-i editing of endogenous transcripts by endog enous adenosine deaminases acting on RNA (ADAR) enzymes, including the ubiquitously and constitutively expressed ADAR1 p110 isoform. We show that fully chemically modified AlMers with chimeric backbones containing stereopure phosphorothio ate and nitrogen-containing linkages based on phosphoryl guanidine enhanced potency and editing efficiency 100-fold com pared with those with uniformly phosphorothioate-modified backbones in vitro. In vivo, AlMers targeted to hepatocytes with N-acetylalactosamine achieve up to 50% editing with no bystander editing of the endogenous ACTB transcript in non-human primate liver, with editing persisting for at least one month. These results support further investigation of the therapeutic potential of stereopure AlMers.

Recruiting endogenous RNA-editing enzymes using chemi vehicles, such as viral vectors or lipid nanoparticles, for application cally modified oligonucleotides holds promise for treating beyond cell culture So far, these technologies have yielded nominal human disease. The most common mutation in human genes editing in vivo®.

is transition from cytosine (C) to thymine (T)<sup>1</sup>, and CpG dinucleo tides are well established hot spots for disease-causing mutations oped relatively short oligonucleotides that elicit A-to-IRNA editing The ADAR family of enzymes catalyze adenine (A)-to-inosine (I) with high efficiency using endogenous ADAR enzymes. These oligo channes in the transcritotme<sup>2</sup> Recause I is read as cuanine (G) nucleotides called AlMers are short and fully chemically modified

- Specificity in vitro & in vivo (NHPs)
- In vitro-in vivo translation (NHPs)
- GalNAc conjugation
- Foundational AIMer SAR



Monian et al., 2022 published online Mar 7, 2022; doi: 10.1038.s41587-022-01225-1 SAR structure-activity relationship

# AIMers: Realizing the potential of therapeutic RNA editing by harnessing endogenous ADAR

Solved for key therapeutic attributes for potential best-in-class RNA editing therapeutics



- Systematized AIMer design enables rapid advancement of new targets
- Strong and broad IP in chemical and backbone modifications, stereochemistry patterns, novel and proprietary nucleosides

# SERPINA1 Z mutation: The most common cause of Alpha-1 antitrypsin deficiency (AATD)



# SERPINA1 Z mutation: The most common cause of Alpha-1 antitrypsin deficiency (AATD)





# RNA editing is uniquely suited to address the therapeutic goals of AATD

### Wave ADAR editing approach addresses all treatment goals:



## ~200K people in US and EU with mutation in *SERPINA1* Z allele (PI\*ZZ)

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only lung manifestations

AAT: Alpha-1 antitrypsin Strnad et al., 2020 N Engl J Med 382:1443-55; Blanco et al., 2017 Int J Chron Obstruct Pulmon Dis 12:561-69; Remih et al., 2021 Curr Opin Pharmacol 59:149-56.

# Robust SERPINA1 RNA editing in mouse model for AATD restores M-AAT protein in serum

RNA editing day 7 AAT serum protein concentration day 7 (3x5 mg/kg, SC)(3x10 mg/kg, SC)60-1000-1000-MZ (50%) editing 800-800-Serum AAT protein (ug/ml) (Mean, s.e.m) 40-Эте 11 µM 600-600-11 µM % SERPINA1 M-AAT (wild-type) 400-400-85% 20-M-AAT (wild-type) 200 200 Z-AAT Z-AAT (mutant) (mutant) PBS SA1-4 SA1-5 SA1-4 PBS SA1-5 PBS



Left: AIMers administered huADAR/SERPINA1 mice (3x5 mg/kg) on days 0, 2, and 4. Livers collected on day 7, and SERPINA1 editing was quantified by Sanger sequencing (shown as mean ± sem) Stats: One-way ANOVA was used to test for differences in editing between SA1-4 and other oligos \* P<0.05. Right: huADAR/SERPINA1 mice administered PBS or 3x10 mg/kg AIMer (days 0, 2, and 4) SC. Proportion of AAT protein in serum measured by mass spec, total AAT protein quantified by ELISA.

## Bi-weekly AIMer treatment results in serum AAT protein levels in mice above anticipated therapeutic threshold

GalNAc-AIMer results in serum AAT protein levels >11 uM at week 19



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SA1-5 administered in huADAR/SERPINA1 mice (8–10 weeks old) Left: Total serum AAT protein quantified by ELISA. Right: Liver biopsies collected at week 19 (one week after last dose); SERPINA1 editing was quantified by Sanger sequencing; Proportion of AAT protein in serum measured by mass spec, total AAT protein quantified by ELISA

## Bi-weekly AIMer treatment supports robust RNA editing and M-AAT protein expression in mice



SA1-5 administered in huADAR/SERPINA1 mice (8–10 weeks old) Left: Total serum AAT protein quantified by ELISA. Right: Liver biopsies collected at week 19 (one week after last dose); SERPINA1 editing was quantified by Sanger sequencing; Proportion of AAT protein in serum measured by mass spec, total AAT protein quantified by ELISA

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# Bi-weekly AIMer treatment results in functional effects in mouse liver at 19 weeks



AIMer (SA1-5) administered in huADAR/SERPINA1 mice (8–10 weeks old) Left: Total serum AAT protein quantified by ELISA. Right: Liver biopsies collected at week 19 (one week after last dose); SERPINA1 editing was quantified by Sanger sequencing; Proportion of AAT protein in serum measured by mass spec, total AAT protein quantified by ELISA; images from liver biopsies stained with PAS-D or AAT-polymer specific antibody

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# AIMer-directed editing is highly specific in mice

RNA editing only detected at PiZ mutation site in SERPINA1 transcript (mouse liver)



**RNA editing across transcriptome** 



Dose 3x10 mg/kg (days 0, 2, 4) SC. Liver biopsies day 7. RNA-seq to quantify on-target SERPINA1 editing, to quantify off-target editing reads mapped to entire mouse genome; plotted circles represent sites with LOD>3 (N=4), SERPINA1 edit site is indicated

# Opportunity for novel and innovative AIMer therapeutics

## **Correct driver mutations with AIMers**



### Restore or correct protein function



Upregulate expression Modify function Modulate protein-protein interaction Post-translational modification Alter folding or processing

### **Examples** AATD

Rett syndrome

Recessive or dominant genetically defined diseases

**Examples** Haploinsufficient diseases Loss of function Neuromuscular Dementias Familial epilepsies Neuropathic pain



## **Modulate protein interactions with AIMers**



- Our RNA editing platform capability allows for correction of driver mutations of AATD
- We have developed RNA editing oligonucleotides AIMers to correct homozygous "ZZ" mutations back to "MZ" heterozygous state
- AIMers lead to durable expression of M-AAT protein and reductions in liver aggregates
- IND-enabling studies for AATD expected to initiate in Q3 2022
- We have established an RNA editing platform for liver and beyond



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#### nature biotechnology

#### **ARTICLES** https://doi.org/10.1038/s41587-022-01225-1

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Technologies that recruit and direct the activity of endogenous RNA-editing enzymes to specific cellular RNAs have therapeutic potential, but translating them from cell culture into animal models has been challenging. Here we describe short, chemically modified oligonucleotides called AlMers that direct efficient and specific A-to-I editing of endogenous transcripts by endog enous adenosine deaminases acting on RNA (ADAR) enzymes, including the ubiguitously and constitutively expressed ADAR1 p110 isoform. We show that fully chemically modified AlMers with chimeric backbones containing stereopure phosphorothio ate and nitrogen-containing linkages based on phosphoryl guanidine enhanced potency and editing efficiency 100-fold com pared with those with uniformly phosphorothioate-modified backbones in vitro. In vivo, AlMers targeted to hepatocytes with N-acetylgalactosamine achieve up to 50% editing with no bystander editing of the endogenous ACTB transcript in non-human primate liver, with editing persisting for at least one month. These results support further investigation of the therapeutic potential of stereopure AlMers.

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is transition from cytosine (C) to thymine (T)<sup>1</sup>, and CpG dinucleo Leveraging our oligonucleotide chemistry platform, we devel tides are well established hot spots for disease-causing mutations oped relatively short oligonucleotides that elicit A-to-I RNA editing The ADAR family of enzymes catalyze adenine (A)-to-inosine (I) with high efficiency using endogenous ADAR enzymes. These oligo changes in the transcriptome?. Because I is read as quanine (G) nucleotides, called AIMers, are short and fully chemically modified by the translational machiner(\*, ADAR-mediated RNA editing has with stereopure phosphorothioate (PS) and nitrogen-containing the potential to revert these disease-causing transitions at the RNA (PN) linkages based on phosphoryl guanidine. In vitro, they level. The potential scope for application of A-to-I editing is large, enhanced potency and A-to-I editing efficiency compared to uni including modulation of polar or charged amino acids, stop codons formly PS-modified AIMers, and in vivo, N-acetylgalactosamine or RNA regulatory sequences<sup>611</sup>, eliciting diverse functional out (GalNAc)-modified AlMers achieved up to 50% editing with no comes (for example, restored protein expression or function) Chemical modifications are known to confer drug-like proper for at least 1 month.

bystander editing in non-human primate (NHP) liver that persisted

ties to oligonucleotides. We set out to determine whether control

over backbone chemistry and stereochemistry and other chemical Results

modifications to an oligonucleotide (Fig1 and Supplementary Note AIMers support RNA editing. To evaluate RNA-editing efficiency 1) can be optimized to elicit sequence-specific A-to-I RNA editing in mammalian cells, we created a luciferase reporter with genes with endogenous ADAR enzymes. As therapeutics, reversible RNA from Gaussia (Gluc) and Cypridinia (Cluc). In the absence of edit editing with oligonucleotides may represent a safer option thaning, only Gluc is expressed, whereas A-to-I editing permits expres those that edit genomic DNA Early technologies designed to elicit sion of Cluc, providing a measure of RNA-editing efficiency and RNA editing in vitro required an exogenous enzyme and an oligo protein expression (Extended Data Fig. 1a). AlMers were designed nucleotide<sup>2-17</sup>. These approaches led to overexpression of editing to mimic naturally occurring double-stranded RNA ADAR subenzyme and substantial off-target editing 416-15, Recent advances strates, as in the GluR2 transcript 413-20 (Extended Data Fig. 1b), have overcome the need for exogenous enzymes in vitfo, but To benchmark RNA editing, we transfected 293T cells with the they still use long oligonucleotides that require ancillary delivery reporter and exogenous ADAR enzyme in the presence or absence

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