

The Endosomal Escape Vehicle Platform Enhances the Delivery of Oligonucleotides to Skeletal and Cardiac Muscle

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ENTRADA'S MISSION Treating Devastating Diseases With Intracellular Therapeutics

THE NEED FOR INTRACELLULAR THERAPEUTICS

Approximately 75% of all disease-causing targets are intracellular and difficult to reach, representing a significant issue when developing effective therapies



The Endosomal Escape Vehicle (EEV[™]) Platform aims to solve the fundamental problem: Lack of efficient cellular uptake and escape from the endosome

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Entrada's pipeline includes a diverse array of high potential and high value assets





EEV PLATFORM



ENDOSOMAL ESCAPE VEHICLE (EEV[™]) PLATFORM

Entrada seeks to solve a fundamental problem: lack of efficient cellular uptake and escape from the endosome; Both are critical to intracellular target engagement and therapeutic benefit

- Unique chemistry results in improved uptake and endosomal escape
- Cyclic structure designed to extend half life and increase stability
- Phospholipid binding potentially enables broad biodistribution to all cells
- Mechanism of internalization conserved across species



A BROADLY APPLICABLE PLATFORM



Entrada has demonstrated intracellular uptake of unique moieties ranging from 1 kDa to 600 kDa



FUNCTIONAL DELIVERY FOR TARGET TISSUES

EEV-therapeutic candidates can be designed to enhance functional delivery to target tissues

Discovery Engine for Intracellular Therapeutics



- High-throughput **EEV library screening** in vitro
- Functional validation of lead EEVs with PMO therapeutic modality in vitro and in vivo
- **EEV** and **linker** optimized for the functional delivery to target tissues *in vivo*

Functional Delivery in the EGFP-654 Transgenic Mice¹



PMO, phosphorodiamidate morpholino oligomer; EGFP-654 transgenic mouse model contains an EGFP gene interrupted by human beta-globin intron 2 with mutated nt654 (Sazani, P. et al. *Nature Biotech.* 2002); PMO654, splicing switching PMO targeting nt654; shown as mean ± standard deviation. ¹Li X. et al. *Mol Therapy-Nucl Acids.* 2023.

TRANSLATION FROM UPTAKE TO OUTCOMES Murine Example

EEV-therapeutic candidates have demonstrated favorable pharmacological properties: efficient intracellular delivery, significant uptake in target tissues and potent pharmacodynamic outcomes





DUCHENNE MUSCULAR DYSTROPHY



SIGNIFICANT THERAPEUTIC NEED EXISTS FOR THE TREATMENT OF DMD

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Duchenne is caused by mutations in the DMD gene, which lead to a lack of functional dystrophin, causing progressive loss of muscle function throughout the body

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Exon skipping therapeutics have been approved based on modest improvement in dystrophin levels ranging from ~1 to 6%



REPEAT EEV-PMO TREATMENT RESTORES MUSCLE INTEGRITY IN D2-mdx MICE

Robust exon 23 skipping after four monthly IV doses of EEV-PMO-23 in D2-mdx mice

Broad dystrophin expression and restoration of muscle integrity after four monthly IV doses of EEV-PMO-23 in D2-mdx mice



PMO-23 EEV-PMO-23

n.s.

Vehicle

PMO-23 EEV-PMO-23

20



Representative Immunofluorescence of Gastrocnemius Muscle (Dystrophin Staining in Green)



Representative Histopathology of Gastrocnemius Muscle (H&E Staining)



• D2-mdx mice (male, n=6-7) were treated with 4 monthly doses of either vehicle, 20 mg/kg PMO-23 or 20 mg/kg PMO-23 equivalent of EEV-PMO-23, and the data were collected ~4 weeks after the last dose

> EEV, Endosomal Escape Vehicle; PMO-23, mouse Dmd exon 23 skipping phosphorodiamidate morpholino oligomer; D2-mdx is a DMD mouse model with a nonsense mutation in DMD exon 23 (Colev et al. Hum. Mol. Genet. 2016); ****p<0.0001; n.s., not significant; shown as mean ± standard deviation.

Vehicle

COMPARISON TO ALTERNATIVE R6-PMO



EEV-PMO significantly improved exon 23 skipping after 3 days in *mdx* mice as compared to competitive R6-PMO



^{*}p<0.05, **p<0.01

• EEV-PMO-23 demonstrates significantly improved PD effects after single 40 mg/kg IV dose in *mdx* mice

ENTR-601-44 IN NHP



A single IV dose of ENTR-601-44 resulted in robust exon skipping in both skeletal and cardiac muscles in NHP, as well as prolonged duration of effect for at least 12 weeks



 At 7 days post IV infusion of 30 mg/kg (PMO equivalent), robust exon 44 skipping across different muscle groups isolated from the ENTR-601-44 treated NHP

Duration of Effect in NHP Biceps for at Least 12 Weeks



 Post IV infusion of 35 mg/kg (PMO equivalent), robust exon 44 skipping observed in biceps in the ENTR-601-44 treated NHP (n=3 per cohort) for at least 12 weeks

ENTR-601-45 IN VITRO EFFICACY

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ENTR-601-45 showed robust exon skipping and dystrophin production in patient-derived skeletal and cardiac muscle cells



ENTR-601-45 TARGET ENGAGEMENT IN HUMAN DMD MICE

ENTR-601-45 delivered up to a two-fold improvement in exon skipping when compared to an equivalent dose of the same EEV conjugated to a casimersen sequence



- A single IV dose of ENTR-601-45 resulted in high levels of exon skipping in hDMD in the mouse skeletal muscle and heart after 1 week
- Both EEV-PMO45 (casimersen sequence) and ENTR-601-45 utilize the same EEV
- EEV-PMO45 uses the same oligonucleotide sequence as casimersen

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DMD DATA SUMMARY

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These results demonstrate that the EEV Platform efficiently delivers oligonucleotides to skeletal and cardiac muscles in preclinical models of Duchenne muscular dystrophy

ENTR-601-44^a

- High levels of exon skipping across mdx, D2-mdx, human dystrophin (hDMD) mouse and NHP studies
- Exon skipping translates to promising dystrophin production in heart and skeletal muscles
- Dystrophin production observed to result in functional improvement

Entrada received authorization in the U.K. to initiate a Phase 1 clinical trial in healthy volunteers

• First participant was dosed in September 2023 with data anticipated in 2H 2024

ENTR-601-45

- Robust exon skipping and dystrophin protein production in patient-derived cardiac and skeletal muscle cells
- High levels of exon skipping were measured in hDMD mouse heart and skeletal muscle tissue

Entrada is planning for regulatory submission in Q4 2024

 Planning for a direct to MAD trial in Duchenne patients for ENTR-601-45

ADDITIONAL PLATFORM OPPORTUNITIES



Entrada continues to invest in and build upon our EEV platform to extend our efforts in developing novel EEV-therapeutic candidates

Target		Platform Approach		Goal
	DNA	yr.	Gene editing	Deliver CRISPR enzyme and repair gene function with guide RNA
JULIUM P	RNA	200 - 200 -	RNA editing	Deliver oligonucleotide therapeutics for RNA editing
			RNA splicing	Modify RNA via exon/intron splicing to activate protein expression
			RNA blocking	Block trinucleotide repeats in RNA to inhibit adverse binding
			RNA silencing	Silence or knockdown RNA to prevent protein expression
	Protein	* * }/~	Protein replacement	Replace proteins and enzymes
			Protein inhibition	Inhibit protein signaling pathways
			Protein degradation	Degrade disease-causing proteins

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