



LENZ

THERAPEUTICS

Topline INSIGHT Results

Phase 2C Clinical Trial NCT05294328

October 2022

Both LNZ101 and LNZ100 met all endpoints with highly significant response rates to 10 hours

	1 Hour			10 Hour		
	Vehicle	LNZ100	LNZ101	Vehicle	LNZ100	LNZ101
Primary Percentage of subjects ≥ 3-line improvement and no loss in BCDVA ≥ 5 letters	6%	71% p<0.001	56% p<0.001	4%	37% p<0.001	48% p<0.001
Secondary Percentage of subjects ≥ 2-line improvement and no loss in BCDVA ≥ 5 letters	27%	86% p<0.001	78% p<0.001	12%	55% p<0.001	58% p<0.001

Broad patient population

Mean age 56 yo (46-73)
 Refractive Error -3.25D SE to +1.5D SE

Well tolerated

Improved NV without compromising DV in both low and normal light

INSIGHT trial compared LNZ100 and LNZ101 against vehicle on key variables



LNZ100

1.75% Aceclidine

- Ready to use
- Preservative Free Eye Drop

LNZ101

1.75% Aceclidine + Brimonidine

- Extended duration

Objective

To evaluate the safety and efficacy of LNZ101 compared with LNZ100 and vehicle in the treatment of Presbyopia

Primary Endpoint

Percentage of subjects who achieve a 3-line or greater improvement and no loss in BCDVA \geq 5 letters at 1h

Secondary Endpoint

Percentage of subjects who achieve a 2-line or greater improvement and no loss in BCDVA \geq 5 letters at 1h

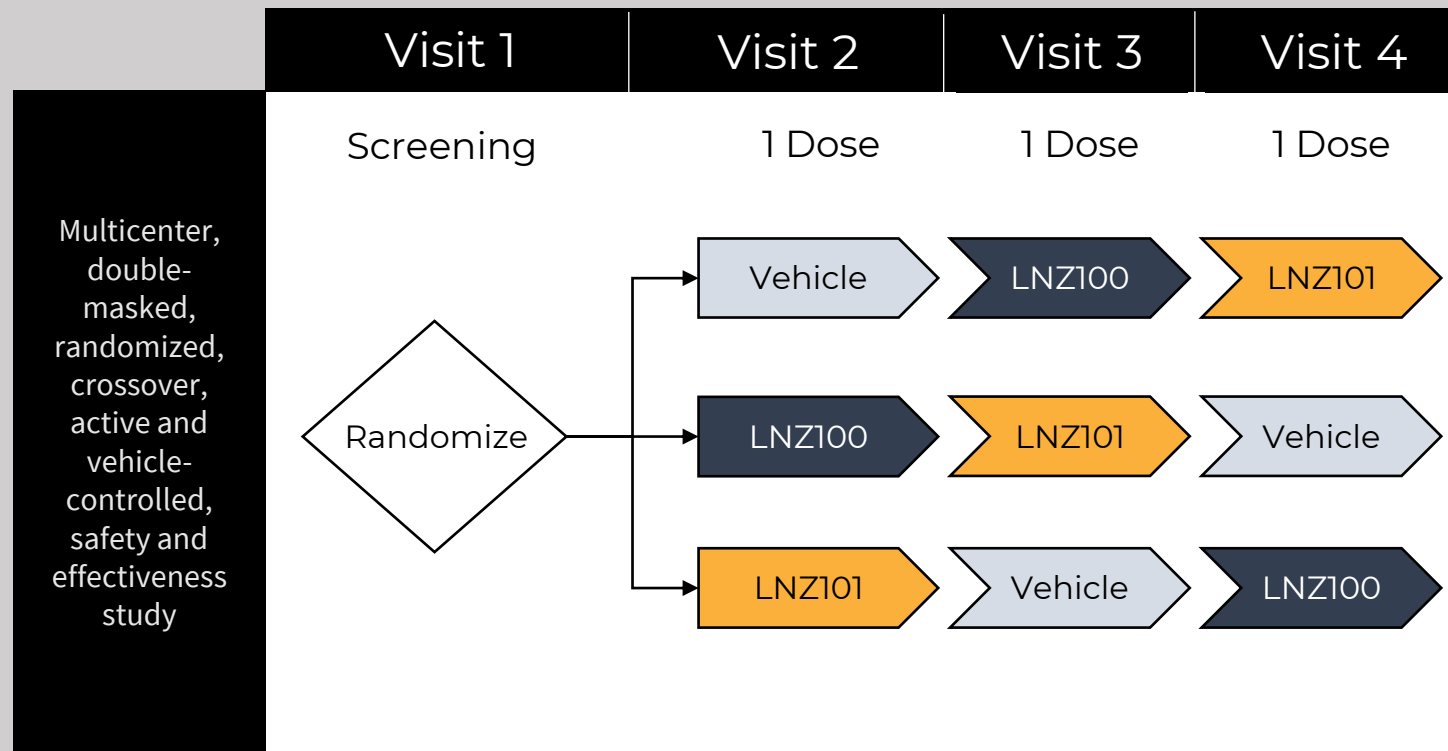
Other Variables

Percentage of subjects who achieved a 3-line or greater improvement no loss in BCDVA \geq 5 letters from 0.5 – 10 hours, Pupil Diameter, AEs

Broad patient population and 10 hour study



INSIGHT Trial Design



Study Design

- 5 US Sites
- 50+ Patients
- Placebo controlled
- 10 hr duration

Study Population

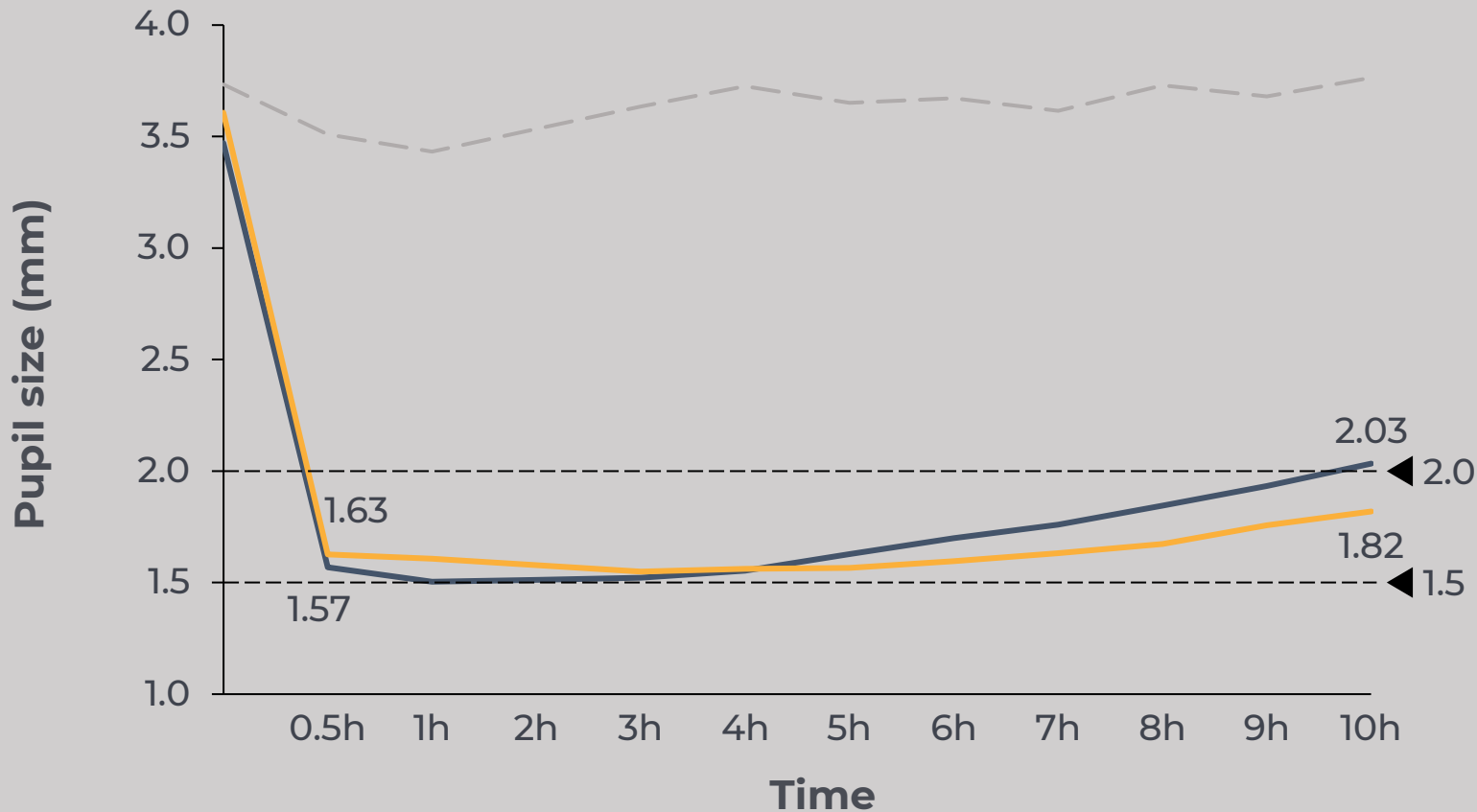
- Average Age: 56 (46 – 73)
- Refractive Rrage (-3.25D SE to +1.5D SE)
- 60%/40% Female/Male
- 60%/40% Brown Iris/Other
- Includes Post Lasik presbyopes and Pseudophakes

Pupil diameter within 1.5mm – 2mm for 10 hours



Pupil Size Near Vision Improvement Biomarker

— LNZ100 (n=49) — LNZ101 (n=50) - - Placebo (n=51)



Average pupil size reduced to **~1.6mm at 30 minutes**

Pupil size **correlates to** lines of **near vision improvement**

Average pupil size **maintained in sweet spot** of 1.5mm to 2mm **for 10 hours**

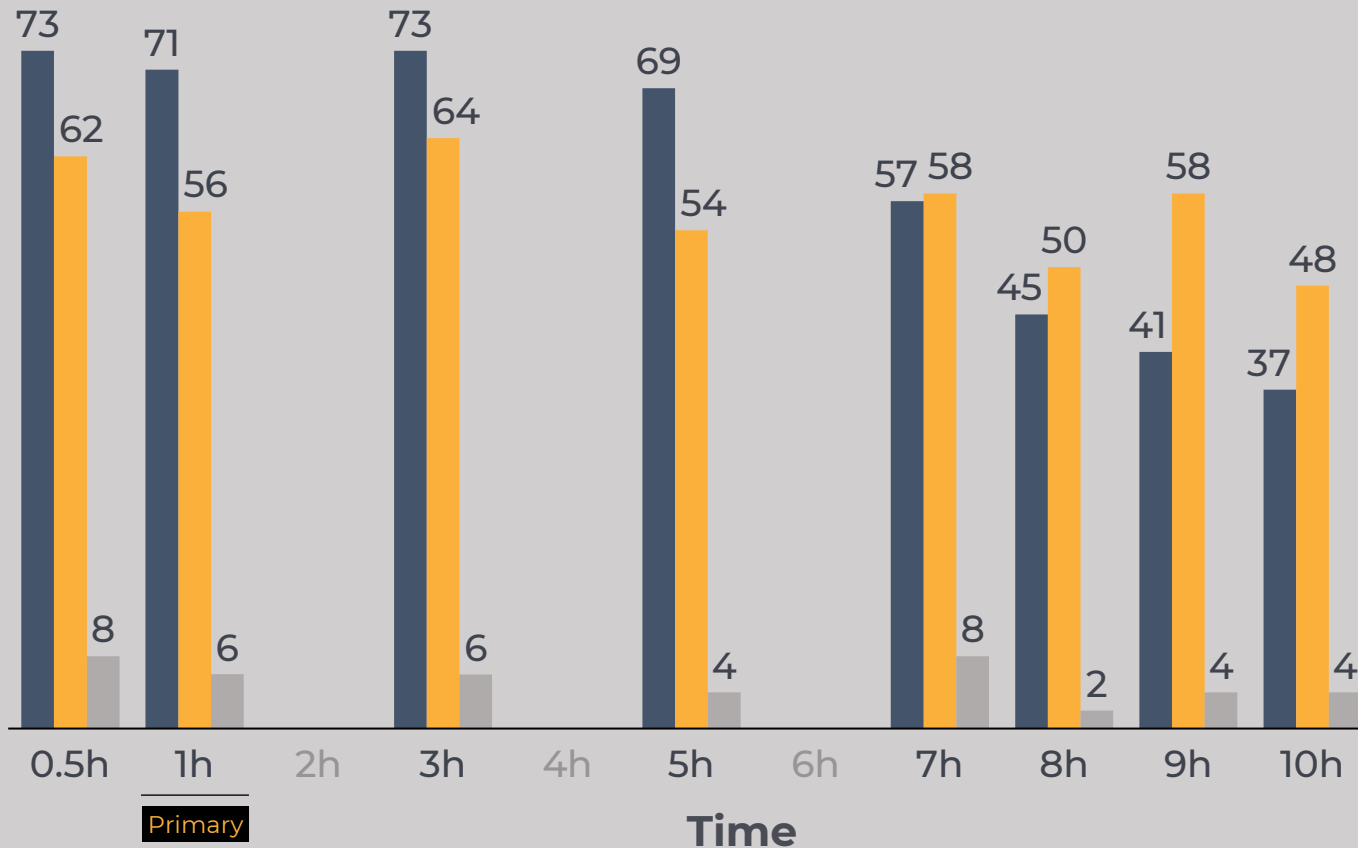
Primary 1 hour endpoint met and 10 hours efficacy

≥3-Line

% Improvement Over Time

(No loss of 5 or more letters distance)

■ LNZ100 (n=49) ■ LNZ101 (n=50) ■ Placebo (n=51)



Extended **category leadership** for efficacy and duration for both LNZ100 and LNZ101

Rapid onset with resp. **73% and 62%** efficacy within **30 min**

Extended Duration with **significance for 10 hours**, LNZ101 statistically separates from LNZ100 at 9 hours

94% of the subjects achieved distance corrected near visual acuity of **20/40 or better**

Well **placebo-controlled** study

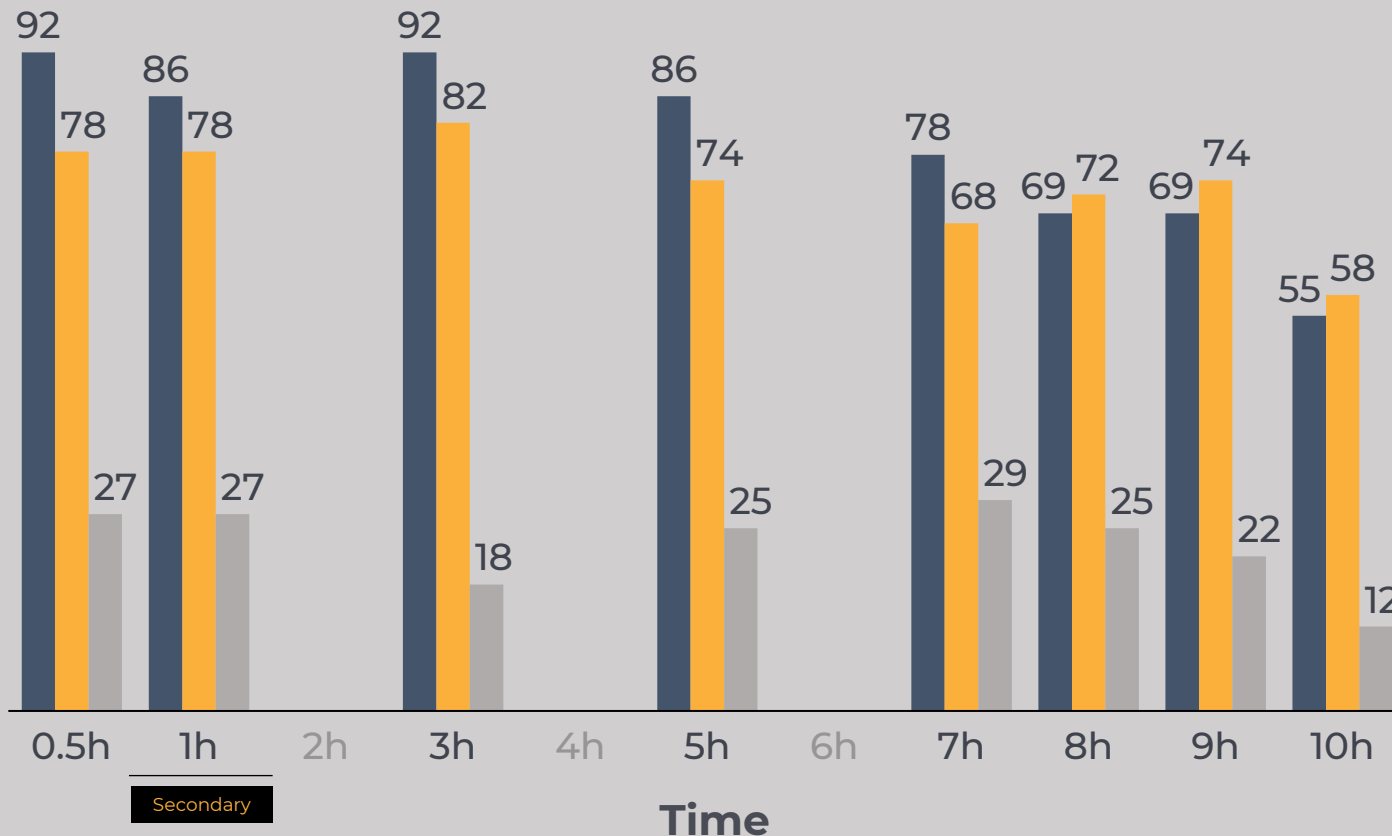
p<0.001 for all time points compared to vehicle

Secondary 1 hour endpoint met and 10 hours efficacy



≥2-Line
% Improvement Over Time
(No loss of 5 or more letters distance)

■ LNZ100 (n=49) ■ LNZ101 (n=50) ■ Placebo (n=51)



Both LNZ100 and LNZ101 **provided clinically meaningful 2 lines** or more NV improvement **for almost all patients**

Rapid onset with resp. **92% and 78%** efficacy within **30 min**

Extended duration with **significance for 10 hours**

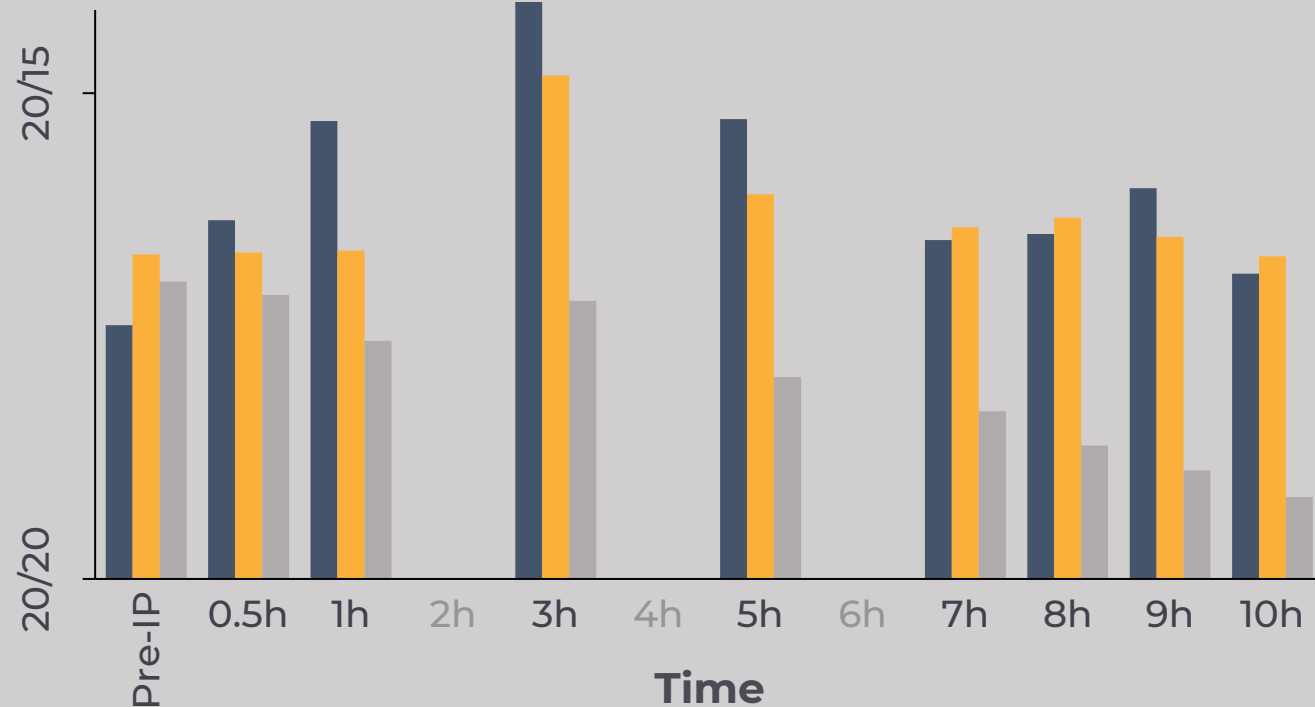
p<0.001 for all time points compared to vehicle

No impact to distance vision in normal and low light



Distance Visual Acuity Best Corrected

■ LNZ100 (n=49) ■ LNZ101 (n=50) ■ Placebo (n=51)

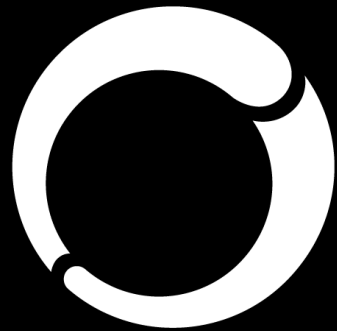


No impact to distance vision in normal light

No impact to distance vision in low light

Well tolerated, No drug related serious adverse events

**Additional INSIGHT data to be provided at
upcoming industry conferences**



LENZ
T H E R A P E U T I C S

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