Kidney Function and Isolated Kidney Transplant Outcomes in Primary Hyperoxaluria Type 1 Treated With Long-term Lumasiran

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Disclosures

Michael J. Somers: Consultancy fees from Alnylam Pharmaceuticals and Dicerna Pharmaceuticals, and scientific review committee chair for ongoing clinical trial with Novo Nordisk

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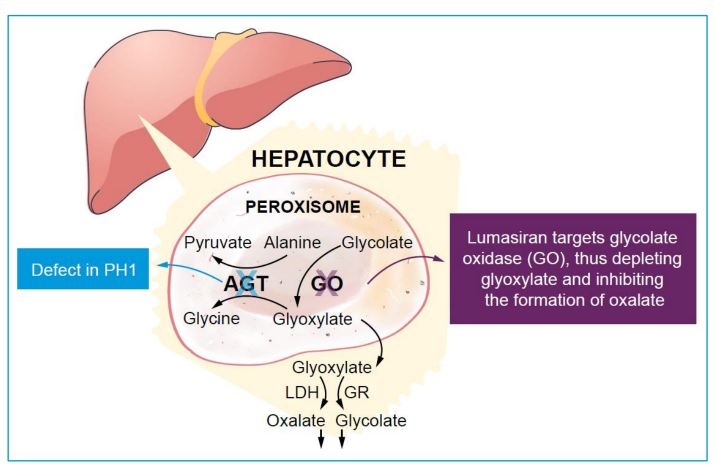
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Introduction

 Primary hyperoxaluria type 1 (PH1) is a rare autosomal recessive disorder associated with hepatic oxalate overproduction, leading to progressive kidney damage



- Historically, PH1 manifests clinically with nephrolithiasis, nephrocalcinosis, and steady decline in eGFR
- Prior to RNA interference therapy with lumasiran, kidney function decline was more rapid as CKD stage advanced
 - Many PH1 patients developed ESKD and required kidney-liver transplantation as treatment
- Lumasiran treatment has demonstrated robust efficacy in reducing urinary oxalate (UOx) and plasma oxalate (POx) and in attenuating kidney injury from PH1, which has spurred reconsideration of expectations for clinical course in CKD and approaches to transplantation in ESKD

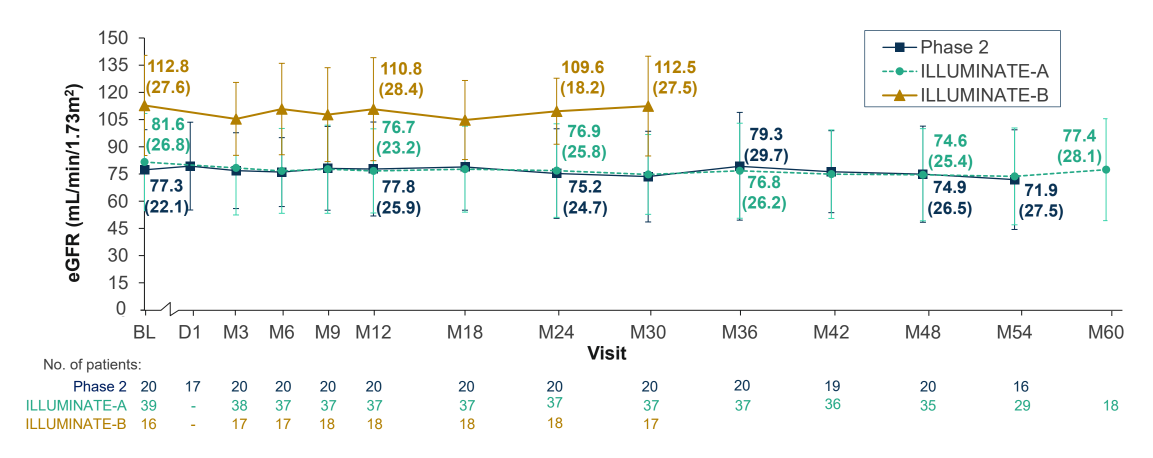
Objective

- In this *post hoc* analysis of lumasiran clinical trials data, we aimed to:
 - o Evaluate rates of eGFR change after lumasiran initiation
 - o Describe post-transplant outcomes following isolated kidney transplant with ongoing lumasiran therapy

Lumasiran Clinical Trials

	Phase 2	ILLUMINATE-A	ILLUMINATE-B	ILLUMINATE-C
Design	Open-label extension study including patients from the single-blind, placebo-controlled Phase 1/2 trial, Part B (in PH1)	Phase 3, randomized, double-blind, placebo- controlled study with extension period	Phase 3, single-arm, open-label study with extension period	Phase 3, single-arm, 2-cohort, open-label study with extension period
Patients, N	20	39	18	21 (6 in Cohort A, 15 in Cohort B)
Inclusion criteria	 PH1 Age 6-64 years eGFR >45 mL/min/1.73m² 24-hour UOx excretion >0.7 mmol/24h/1.73m² Completed Phase 1/2 study, Part B 	 PH1 Age ≥6 years eGFR ≥30 mL/min/1.73m² 24-hour UOx excretion >0.7 mmol/24h/1.73m² 	 PH1 Age <6 years eGFR >45 mL/min/1.73m² if age ≥12 months or normal serum creatinine if age <12 months UOx:Cr greater than ULN for age 	 PH1 All ages eligible eGFR ≤45 mL/min/1.73m² if age ≥12 months or elevated serum creatinine if age <12 months POx ≥20 µmol/L Not on HD at study start (Cohort A); stable HD regimen (Cohort B)
Primary endpoint	Incidence of AEs	% change in 24-hour BSA-corrected UOx at 6 months	% change in UOx at 6 months	% change in POx at 6 months (in Cohort B, % change in pre-dialysis POx)
Total duration	Up to 54 months	Up to 60 months	Up to 60 months	Up to 60 months

eGFR Over Time in Phase 2, ILLUMINATE-A, and ILLUMINATE-B



- The range of baseline eGFR values was 32 to 174 mL/min/1.73m² among patients enrolled in these 3 trials
- Median (range) ages at baseline were 11.5 (6-43) years in Phase 2, 14.0 (6-60) years in ILLUMINATE-A, and 4.2 (0.3-6) years in ILLUMINATE-B

Change in eGFR per Year (slope) in Lumasiran Clinical Trials

• Change in eGFR (slope) ranged from -0.6 to 0.4 mL/min/1.73m2 per year up to 60 months



Rates of eGFR Change in ILLUMINATE-C

- In ILLUMINATE-C, the 6 Cohort A patients were pre-dialysis with eGFR <45 mL/min/1.73m² at study initiation
 - Three patients (study initiation eGFRs all <16.5 mL/min/1.73m²) progressed to need to start dialysis
 - In 2 of 3 patients remaining off dialysis with baseline eGFR data, a relatively modest annualized decline in eGFR was seen through Month 36 while on lumasiran therapy

Cohort A: Patients Remaining Off Dialysis at Month 36

Patient	Baseline eGFR	Month 36 eGFR	Annualized eGFR Decline	
1	24.0	17.2	-2.3	
2	34.1	31.3	-0.9	

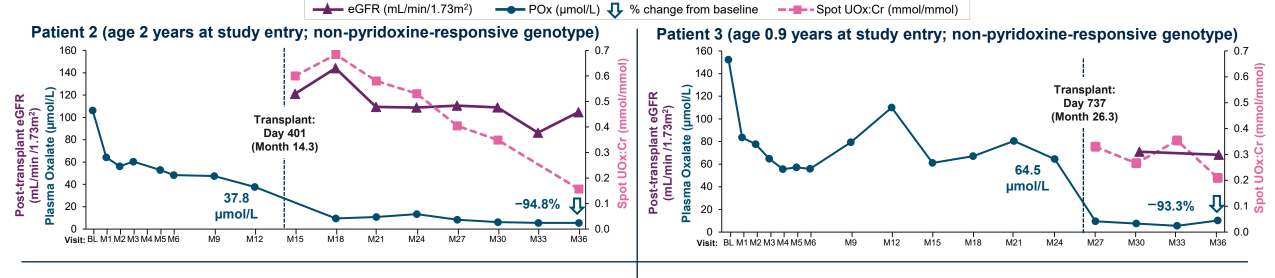
Isolated Kidney Transplant Outcomes in ILLUMINATE-C

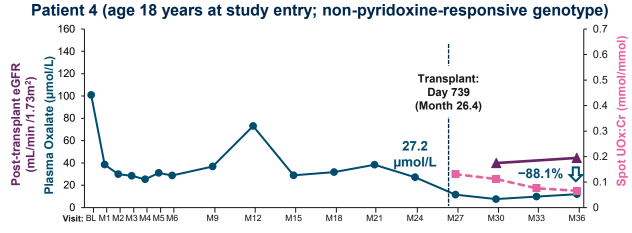
Cohort B: Kidney Transplant by Month 36

Patient	Age at Baseline (Years)	POx at Baseline (µmol/L)	Months of Lumasiran Treatment Until Transplant	POx Pre- transplant (μmol/L)	Months of Post- transplant Follow-up to Month 36	Overall % Decline in POx from Baseline
1	44	84.6	6.7	47.1	29.3	-76.1%
2	2	106.3	14.3	37.8	21.7	-94.8%
3	0.9	152.3	26.3	64.5	9.7	-93.3%
4	18	100.9	26.4	27.2	9.6	-88.1%
5	1	103.7	32.5	19.2	3.6	-93.4%

- Of 15 patients enrolled in Cohort B (on HD at study start), 5 underwent isolated kidney transplant as of Month 36
- All 5 patients had reductions in POx from baseline prior to transplantation (range of POx reduction: -37.5 to -87.8 μmol/L)
 - POx at baseline, range: 84.6 to 152.3 μmol/L; POx at last visit prior to iKT, range: 19.2 to 64.5 μmol/L
- Further reductions post-transplant indicate improved POx clearance with functioning kidney grafts (POx at Month 36, range: 5.6 [LLOQ] to 20.2 μmol/L)

Pediatric Isolated Kidney Transplant Outcomes in ILLUMINATE-C





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- Post-isolated kidney transplant with ongoing lumasiran therapy, POx declined to low levels and remained in a range without systemic risk of oxalate deposition
- UOx levels declined over time post-transplant, reflecting improved oxalate homeostasis and systemic oxalate burden

- eGFR remained satisfactory and in expected post-transplant range
- Reported AEs post-transplant were all unrelated to lumasiran and reflected typical post-transplant clinical events

Patient 5 (age 1 year at study entry; pyridoxine-responsive genotype)

 Results in Patient 1 (age 44 at study entry) were generally consistent with those in pediatric patients

Conclusions



Annual eGFR decline over 30 to 60 months was minimal in patients with PH1 treated with lumasiran in clinical trials encompassing a range of baseline kidney function and age groups (baseline age range: 3 months to 60 years)



Lumasiran treatment effectively lowered POx to allow for consideration of isolated kidney transplantation with end-stage kidney disease



Post-transplant adverse events were frequent and included transplant-related complications (not related to study drug), highlighting the risks associated with organ transplantation



All patients who had isolated kidney transplantation remained dialysis-free, had no oxalate nephropathy, and continued lumasiran treatment post-transplant as of the Month 36 data cut in ILLUMINATE-C

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Thank you to the patients, their families, investigators, study staff, and collaborators for their participation in the lumasiran clinical studies