Study ALLN-177-206: Case Series of First Four Subjects with EH and Advanced CKD Treated with Reloxaliase

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Background

- Enteric hyperoxaluria (EH), a serious metabolic disorder, results from fat malabsorption due to GI surgery and other GI conditions. Despite being a major risk factor for kidney stones it also can lead in severe cases to CKD and ESRD
- Oxalate nephropathy secondary to EH
- No effective pharmacological treatment exists and dietary restriction, calcium supplementation, and high liquid intake is the only therapeutic approach. Difficult to sustain and/or suboptimal effect
- Reloxaliase is an oral enzyme, oxalate-specific therapy that degrades oxalate in the GI tract
- This basket study is enrolling patients ≥12 years of age to examine the potential of reloxaliase to reduce both UOx and POx

Hypothesis

- Declining kidney function leads to oxalate accumulation in plasma (hyperoxalemia) and body (systemic oxalosis)
- With declining kidney function, the GI tract has the potential to play a more significant role in reduction of oxalate burden
- By degrading oxalate in the GI tract, reloxaliase may be able to reduce oxalate burden in the body measured by UOx and POx reduction

Open-Label 3-Month Basket Study

Study Objectives:
Evaluate reloxaliase efficacy and tolerability in subjects with severe enteric hyperoxaluria (EH with CKD/ESRD and hyperoxalemia)

End Point:
Change from baseline in plasma oxalate (POx) and urine oxalate (UOx/d)

Rationale:
- Basket study in severe forms of hyperoxaluria: EH with CKD, dialysis or post kidney transplant with hyperoxalemia, PH1-3
- Pilot-testing use of plasma oxalate (POx) in a multicenter trial for potential future studies
- First exposures in dialysis, PH, and adolescents

Screening and Baseline: Up to 4 Weeks

<table>
<thead>
<tr>
<th>Key Entry Criteria</th>
<th>Treatment Period: 3 Months</th>
<th>Follow-Up: 4 Weeks</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Week 4</td>
<td>Week 8</td>
</tr>
<tr>
<td></td>
<td>Reloxaliase 7,500 u with meal/snacks 5xd for 12 weeks, up to 37,500 u/d</td>
<td>1 x POx 2 x UOx/d</td>
</tr>
<tr>
<td></td>
<td>1 x POx 2 x UOx/d</td>
<td>1 x POx 2 x UOx/d</td>
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<tr>
<td></td>
<td>2 x POx 3 x UOx/d</td>
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</tbody>
</table>

Self-selected diet recorded in diaries; blood samples collected in the morning after an overnight fast or prior to new dialysis sessions

Urine and Plasma Oxalate Reduction in Subjects with CKD

<table>
<thead>
<tr>
<th>Short Bowel Syndrome</th>
<th>Fat Malabsorption</th>
<th>Crohn’s Disease</th>
<th>Chronic Pancreatitis</th>
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</thead>
<tbody>
<tr>
<td>CKD 3b, BMI 23.1</td>
<td>CKD 3t, BMI 19.7</td>
<td>Hemodialysis 2y, BMI 26.4</td>
<td>Hemodialysis 2y, BMI 19.2</td>
</tr>
<tr>
<td>EH history 6y M, 76y</td>
<td>KTx 2015, oxalosis 3 mo after F, 72y</td>
<td>M, 69y</td>
<td>M, 61y</td>
</tr>
<tr>
<td>UOx: 98 mg/g</td>
<td>UOx: 210 mg/g</td>
<td>POx: 40.4 µmol/L</td>
<td>POx: 104.2 µmol/L</td>
</tr>
<tr>
<td>POx: 6.8 µmol/L</td>
<td>POx: 9.0 µmol/L</td>
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</tbody>
</table>

- Mean % Change in UOx and POx

Summary

- Reloxaliase was well tolerated and reduced both UOx and POx, suggesting the potential for reducing systemic oxalate deposition with chronic therapy
- Reduction in plasma oxalate, if sustained, can potentially delay need for dialysis, reduce the frequency of dialysis sessions, help qualify for kidney transplant or protect a new kidney post-transplant
- These initial data support further testing of reloxaliase in patients with severe EH and CKD with hyperoxalemia

Acknowledgment: This research was sponsored by Allena Pharmaceuticals, and was made possible by the valuable contributions of participating investigators and patients. We want to thank all patients and families for participating, this study is ongoing and please contact us if you are interested in taking part in the study.

References:
1) Lumlertgul et al KI 2018
2) Woodcock et al NIM 2017. This poster does not present data for PH.