Extrapolation & Pediatric Development: A case study from pediatric Ulcerative Colitis

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Agenda

• Rationale for extrapolation in pediatric development
• Justification for extrapolation in pediatric UC
• Golimumab pediatric UC case study
• Key outstanding questions
Rationale for Extrapolation in Pediatric Development

- Timely access to approved treatments in Pediatric patients!
- Take advantage of development and approval process for adult indications
- Avoids limitations of pediatric efficacy studies
  - Difficulty of enrolling placebo-controlled trials
  - Limitations of small, underpowered studies
- Avoidance of unnecessary clinical studies in pediatrics
Justification for Extrapolation in Pediatric UC

Disease course is similar in children and adults

Treatment effects are similar in children and adults

Exposure-response relationship is comparable in children and adults
Similarity of Disease

• Pathogenesis and genetics of UC similar in adults and children

• “Although some differences in disease severity exist, the pathogenesis of UC in adults and children is the same, and the disease course in these two populations are similar enough to allow extrapolation of efficacy outcomes from adults to children*.”

• However, children tend to have more “severe disease” as evidenced by a higher incidence of pan-colonic disease

*Mulberg, JPGN 2014
Similarity of Treatment Effects

Across medications, treatment effects are generally similar between children and adults

<table>
<thead>
<tr>
<th></th>
<th>Adults</th>
<th>Pediatric</th>
</tr>
</thead>
<tbody>
<tr>
<td>5-ASAs</td>
<td>50%</td>
<td>59-70%</td>
</tr>
<tr>
<td>Corticosteroids</td>
<td>60%</td>
<td>58%</td>
</tr>
<tr>
<td>6-MP/AZA</td>
<td>49%</td>
<td>53%</td>
</tr>
<tr>
<td>TNFs (infliximab)</td>
<td>69%</td>
<td>73%</td>
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</table>

Similarity of Exposure-Response

- Similarity of Exposure Response in Pediatric and Adult UC has been previously demonstrated with anti-TNF therapy (ie. Infliximab)

![Graph showing similarity of exposure-response in pediatric and adult UC](image)

Agenda

• Rationale for extrapolation in pediatric development
• Justification for extrapolation in pediatric UC
• Golimumab pediatric UC case study
• Key issues & group discussion
Golimumab Peds UC Case Study

- Molecular analysis of Pediatric UC to confirm similarity

- **Initial PK study across full pediatric age range**
  - Molecular analyses, PK, efficacy, E-R, Safety in children
  - Compare results to adult program to address “similarity” requirements

- **Modeling & Simulation analyses**
  - Further evaluate similarity of PK and E-R between children and adults

- **Goal:** Find the right dose of golimumab to safely and effectively treat pediatric ulcerative colitis
Molecular Analyses to Support Demonstration of Similarity of Disease

- Similarity in molecular response to golimumab in adults and children was also demonstrated

- Similarity in molecular profile of limited and extensive disease for both adult and pediatric UC was also demonstrated
CNTO148UCO1001 Study Overview

- Study consists of a PK portion (Week 0-14) and a study extension (Week 14-126); PK and E-R data through Week 14 are reported here

- Patients were dosed based on body weight at baseline:
  - <45 kg: 90 mg/m² at Week 0; 45 mg/m² at Week 2 and q4w in responders
  - ≥45 kg: 200 mg at Week 0; 100 mg at Week 2 and q4w in responders

- Blood samples were collected through Week 14 to evaluate serum golimumab concentrations and immunogenicity

- Efficacy outcomes were assessed using the Mayo score and Pediatric Ulcerative Colitis Activity Index (PUCAI) at Week 6

![Timeline Diagram]
Rates of Clinical Response, Remission, and Mucosal Healing compared in adults and children

*Similar results by age and weight (dose regimen) subgroups in the paediatric study
Consistent Outcomes by Pediatric Subgroups

Example shown: Mayo Clinical Response at Week 6

- Pediatric (CNT0148UCO1001)
  - N = 10, 2 to 11 years: 70%
  - N = 25, 12 to 17 years: 56%
  - N = 15, Golimumab 90 mg/m²: 67%
  - N = 20, Golimumab 200 mg: 55%
  - N = 35, Total: 60%

- Adult (C0524T17)
  - N = 257, Golimumab 200 mg: 52%
  - N = 256, Placebo 100 mg: 30%
### Similar PK (Descriptive Analyses)

<table>
<thead>
<tr>
<th>Visit</th>
<th>Mean (±SD) golimumab concentration (µg/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Week 2</td>
<td>6 (±2) Pediatric, 6 (±2) Adult</td>
</tr>
<tr>
<td>Week 4</td>
<td>4 (±2) Pediatric, 4 (±2) Adult</td>
</tr>
<tr>
<td>Week 6</td>
<td>2 (±2) Pediatric, 2 (±2) Adult</td>
</tr>
<tr>
<td>Week 14</td>
<td>1 (±1) Pediatric, 1 (±1) Adult</td>
</tr>
</tbody>
</table>

**Serum Golimumab Concentrations (µg/ml) at Week 6**

- **Pediatric patients <45 kg**
- **Pediatric patients ≥ 45 kg**
- **Adults**
Exposure-Response (Descriptive Analyses)

Serum golimumab concentrations were positively associated with efficacy outcomes; this relationship was generally comparable between pediatric and adult patients.

**Relationship Between Serum Golimumab Concentrations at Week 6 and Clinical Efficacy Outcomes at Week 6 in the Adult and Pediatric Ulcerative Colitis Populations**
E-R for mucosal healing was similar to that for clinical response

A greater proportion of children achieved clinical remission than adults at the same concentrations
Overall study results were similar to that observed with Infliximab in Pediatric UC

- Clinical Response Mayo Remission PUCAI Remission Mucosal Healing (0/1)

Proportion (%) of Subjects

- Golimumab Peds (UCO1001)
- Infliximab Peds (T72)

Key Outstanding Questions
How is Similarity Of E-R Relationships between children and adults defined?

Clinical Response

Clinical Remission

• What is the benchmark for defining “similar”? Is this descriptive? Or Statistical?

• Is the goal to match the adult exposures and adult response? Or to optimize response?
  – If the remission rates in children are higher than in adults, does that undermine the argument that E-R is “similar”?
How should maintenance data be analyzed?

- Based upon adult data, approximately 50-60% of patients will achieve clinical response, of which 50-60% will maintain response over 1 year → 25-35% of original sample size

  - **PK approach**: PK data from GLM peds UC study demonstrates comparable steady state concentrations through week 14 in children are similar to adults, suggesting target maintenance exposures are achieved in children with UC

  - **Descriptive approach**: Show that maintenance data is consistent with the adult data. This was used for infliximab pediatric UC program

  - **E-R approach**: Demonstrate similar E-R in multiple phases of a disease. This approach will require identical design in maintenance (including additional endoscopies, similar discontinuation criteria, etc...).

  - **Efficacy approach**: Extremely large number of subjects (e.g. > 600) are needed to appropriately power a randomized withdrawal study

- If similar exposure-response is demonstrated in induction, is the bar for maintenance different?
How much safety data is needed?

- 35 children were studied in the UCO1001 study
- Limited safety beyond 14 weeks (ie. 14 subjects beyond 1 year)

However:
- Data was supplemented with 173 children with JIA treated with golimumab
- Large adult database show that the safety of golimumab is similar to other TNFs
- Safety concerns of TNFs are characterized in children
- The most serious events are rare and unlikely to be observed in a pediatric clinical trial.
• What is the size of safety database necessary for approval?
  – Is there a magic number?
  – Are the requirements the same for different diseases/ indications (e.g. UC vs JIA)?

• How should safety from children exposed to the same compound but different indication (e.g. JIA) be factored in?

• Are the safety requirements different for novel MOAs and known MOAs (e.g. anti-TNFs)?
How do we extrapolate from one anti-TNF to another?

- What are the criteria and conditions that need to be met?
- What is the right balance of benefit and risk?

### What is the burden of proof needed to meet the expectations of extrapolation?

<table>
<thead>
<tr>
<th>Key Component</th>
<th>Approach</th>
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<tbody>
<tr>
<td>Rationale for extrapolation</td>
<td>Consensus that new ways to approach pediatric studies is essential</td>
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<tr>
<td>Similar Disease, Similar Response to Treatment</td>
<td>Built upon literature with molecular analysis</td>
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<tr>
<td>Characterize PK in children with Ulcerative Colitis</td>
<td>Conducted a rigorous phase 1 study across the age range and pop PK analysis</td>
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<tr>
<td>Characterize Exposure-Response</td>
<td>Leveraged experience with infliximab in pediatric UC and demonstrate consistent data with golimumab in UC</td>
</tr>
<tr>
<td>Characterize safety of golimumab in children</td>
<td>Large database of anti-TNF use in children and utilize data from golimumab from a different pediatric population</td>
</tr>
<tr>
<td>Identify uncertainties</td>
<td>Limitations of E-R model, limited maintenance data, low number of exposed patients with pediatric UC</td>
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*This is a work in progress for industry, academics, and health authorities*
Goal of Extrapolation: Close the gap between adult and pediatric approvals

<table>
<thead>
<tr>
<th>Compound</th>
<th>Indication</th>
<th>Adult Approval</th>
<th>Pediatric Approval</th>
<th>Gap</th>
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</thead>
<tbody>
<tr>
<td>Infliximab</td>
<td>Crohn’s Disease</td>
<td>1998</td>
<td>2006</td>
<td>8 years</td>
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<tr>
<td></td>
<td>Ulcerative Colitis</td>
<td>2005</td>
<td>2011</td>
<td>6 years</td>
</tr>
<tr>
<td>Adalimumab</td>
<td>Crohn’s Disease</td>
<td>2007</td>
<td>2014</td>
<td>7 years</td>
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<table>
<thead>
<tr>
<th>Pediatric Regulations</th>
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<tr>
<td>Ulcerative Colitis</td>
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<tr>
<td>Golimumab</td>
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*Approximately 15% enrolled in 2 years

Are we closing the gap with the current extrapolation framework?