



AAAAI American Academy of  
Allergy Asthma & Immunology  
**ANNUAL MEETING**  
PHILADELPHIA, PA · MARCH 13-16, 2020

# Clinical Activity of Antolimab (AK002), an Anti-Siglec-8 Monoclonal Antibody, in Treatment-Refractory Chronic Urticaria

Sabine Altrichter<sup>1</sup>; Petra Staubach<sup>2</sup>; Malika Pasha<sup>3</sup>; Henrik S. Rasmussen<sup>3</sup>; Bhupinder Singh<sup>3</sup>; Alan T. Chang<sup>3</sup>; Jonathan A. Bernstein<sup>4</sup>, Frank Siebenhaar<sup>1</sup>, Marcus Maurer<sup>1</sup>

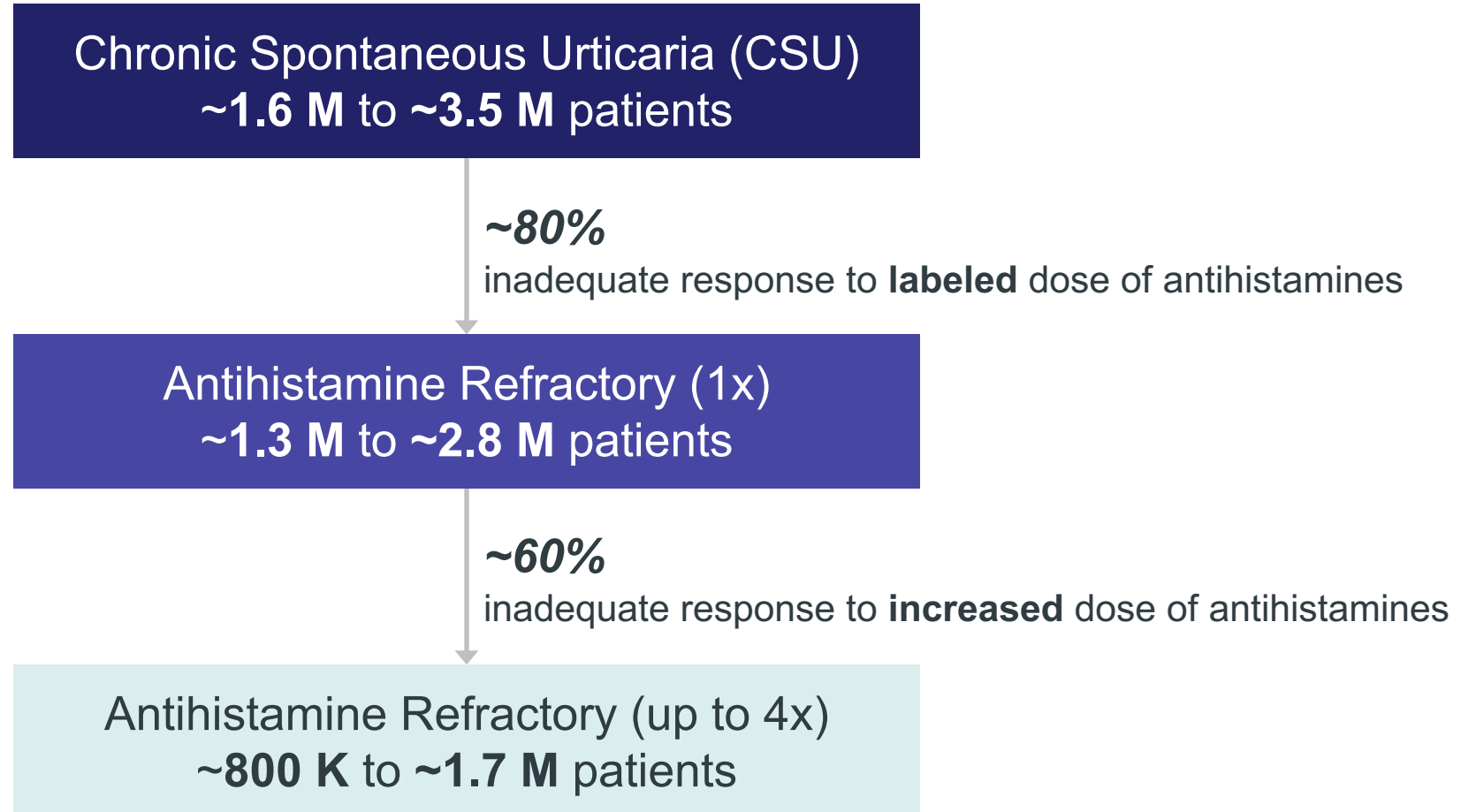
- 1. Charité-Universitätsmedizin Berlin, Berlin, Germany
- 2. University Medical Center Mainz, Mainz, Germany
- 3. Allakos, Inc., Redwood City, CA, United States
- 4. University of Cincinnati, Cincinnati, United States



# Disclosures

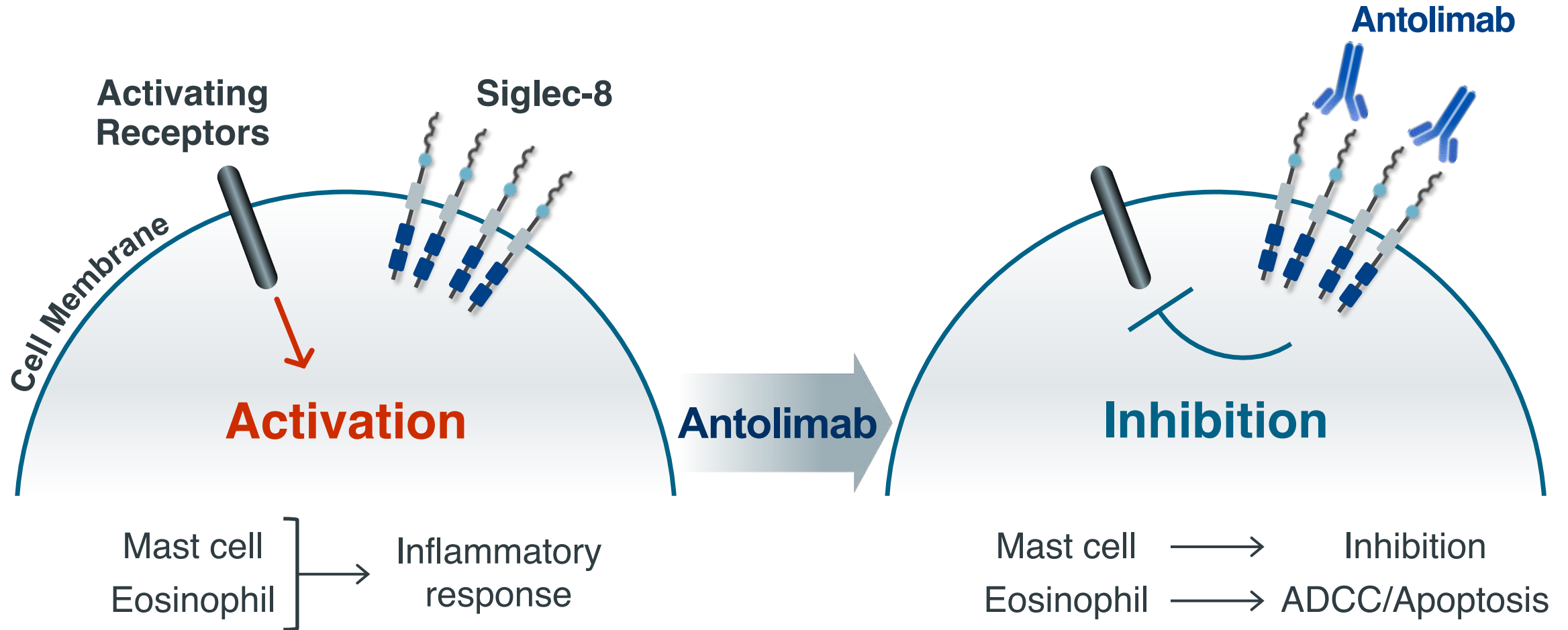
- Sabine Altrichter is an investigator for this study
- Antolimab (AK002) is an investigational drug candidate and is not FDA/EMA approved

# Prevalence of Antihistamine-Refractory CSU in US



Source: Maurer et al. *Allergy*. 2011 Mar;66(3):317-30; Staevska et al. *J Allergy Clin Immunol*. 2010 Mar;125(3):676-82; Van den Elzen et al. *Clin Transl Allergy* (2017) 7:4

# Antolimab (AK002) Targets Siglec-8 on Mast Cells and Eosinophils



# Study Design

Design	Key Endpoints	
<ul style="list-style-type: none"> <li>• Open-label</li> <li>• 45 patients – 4 cohorts               <ul style="list-style-type: none"> <li>- Omalizumab-naïve CSU</li> <li>- Omalizumab-refractory<sup>1</sup> CSU</li> <li>- Cholinergic urticaria</li> <li>- Symptomatic Dermographism</li> </ul> </li> <li>• One month screening period</li> <li>• Dosed once monthly for 6 months</li> <li>• 8 weeks safety follow up</li> <li>• 0.3 mg/kg starting AK002 dose; increased to 1.0 mg/kg (dose 2 and 3); if UCT &lt;12, increased to 3.0 mg/kg (dose 4, 5, and 6)</li> </ul>	<b>Primary</b>	<ul style="list-style-type: none"> <li>• Change in Urticaria Control Test (UCT) Week 22 from BL               <ul style="list-style-type: none"> <li>- Complete Response: UCT score <math>\geq 12</math> and <math>\Delta UCT \geq 3</math></li> <li>- Partial Response: <math>\Delta UCT \geq 3</math></li> <li>- No Response: <math>\Delta UCT &lt; 3</math></li> </ul> </li> </ul>
	<b>Secondary</b>	<ul style="list-style-type: none"> <li>• Change in disease activity by UAS7 (CSU only)</li> <li>• Safety and tolerability</li> </ul>

<sup>1</sup> Patients refractory to omalizumab (n=11): AH treatment up to 4x labeled dose, OMA doses up to 600 mg, average OMA treatment duration ~10 months; UCT on OMA: 4.1

# Urticaria Disease Assessment Tools

## URTICARIA CONTROL TEST (UCT)

- Measures disease control (symptoms and quality of life)
- Used in clinical practice
- Can be used in both chronic spontaneous urticaria and chronic inducible urticarias (e.g. cholinergic and symptomatic dermographism)

## URTICARIA ACTIVITY SCORE (UAS)

- Comprised of measures of itch & hives:
  - Itch Severity Score (ISS)
  - Hives Severity Score (HSS)
- UAS7 used as primary endpoint for regulatory approval in CSU (can only be used in CSU)

**UCT and UAS7 recommended by EAACI CU Guidelines to assess CSU disease**

# Baseline Patient Characteristics

	Chronic Spontaneous Urticaria		Chronic Inducible Urticaria	
	Omalizumab Naïve (n=13)	Omalizumab Refractory (n=11)	Cholinergic Urticaria (n=11)	Symptomatic Dermographism (n=10)
Age	65 (30-75)	29 (22-60)	33 (18-62)	27 (19-56)
Female	100%	82%	55%	60%
BMI	32 (20-44)	26 (20-42)	27 (23-39)	30 (22-36)
UCT	3.2	3.7	5.4	5.7
UAS7	18.5 (8.6-41.0)	28.7 (18.1-35.8)	-	-

# High Response Rate by UCT with Antolimab in CSU

## Chronic Spontaneous Urticaria<sup>1</sup>

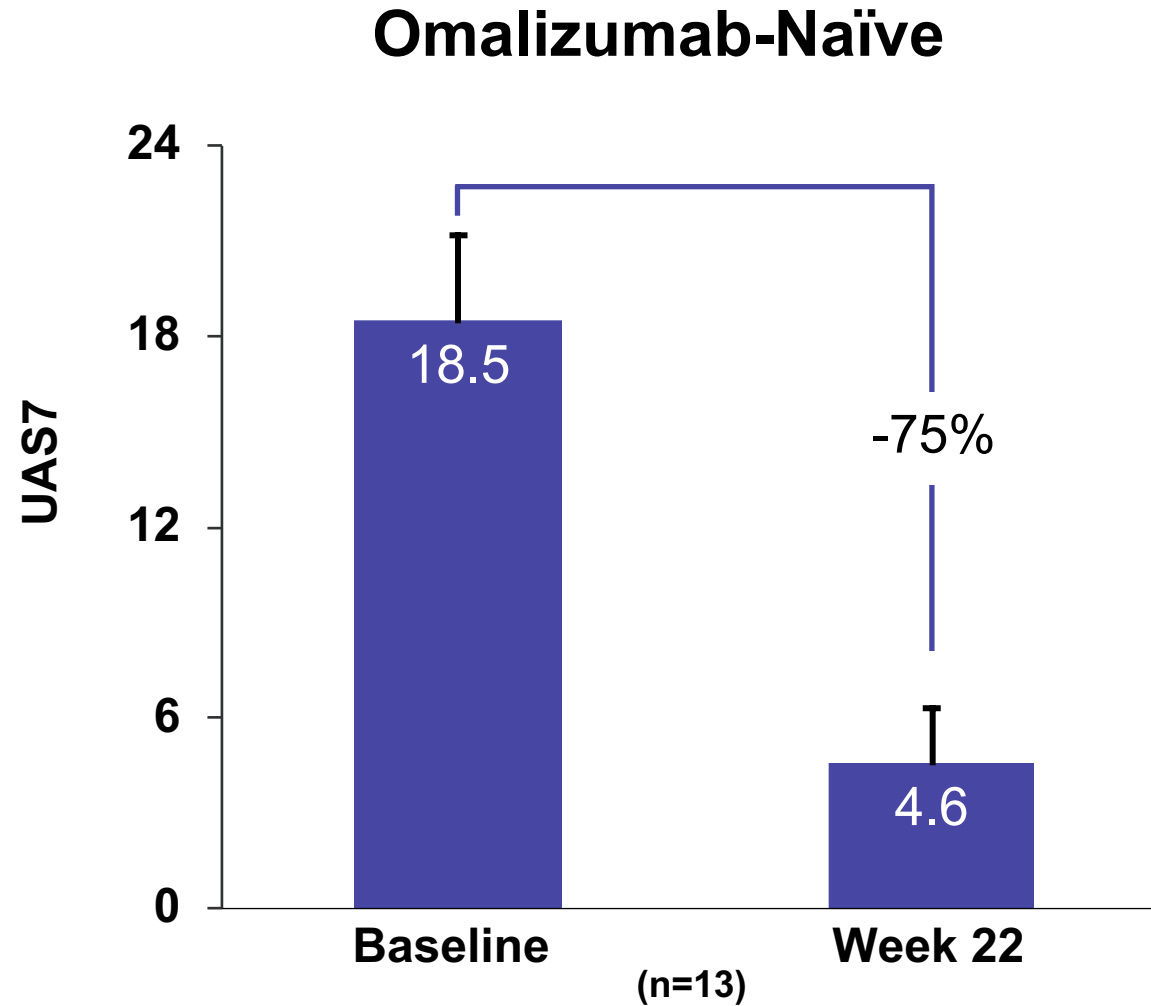
Urticaria Control Test Response <sup>1</sup>	Omalizumab Naïve (n=13)	Omalizumab Refractory (n=7)
Complete Response	92%	57%
Partial Response	0%	29%
No Response	8%	14%

<sup>1</sup> Patients who received all 6 doses

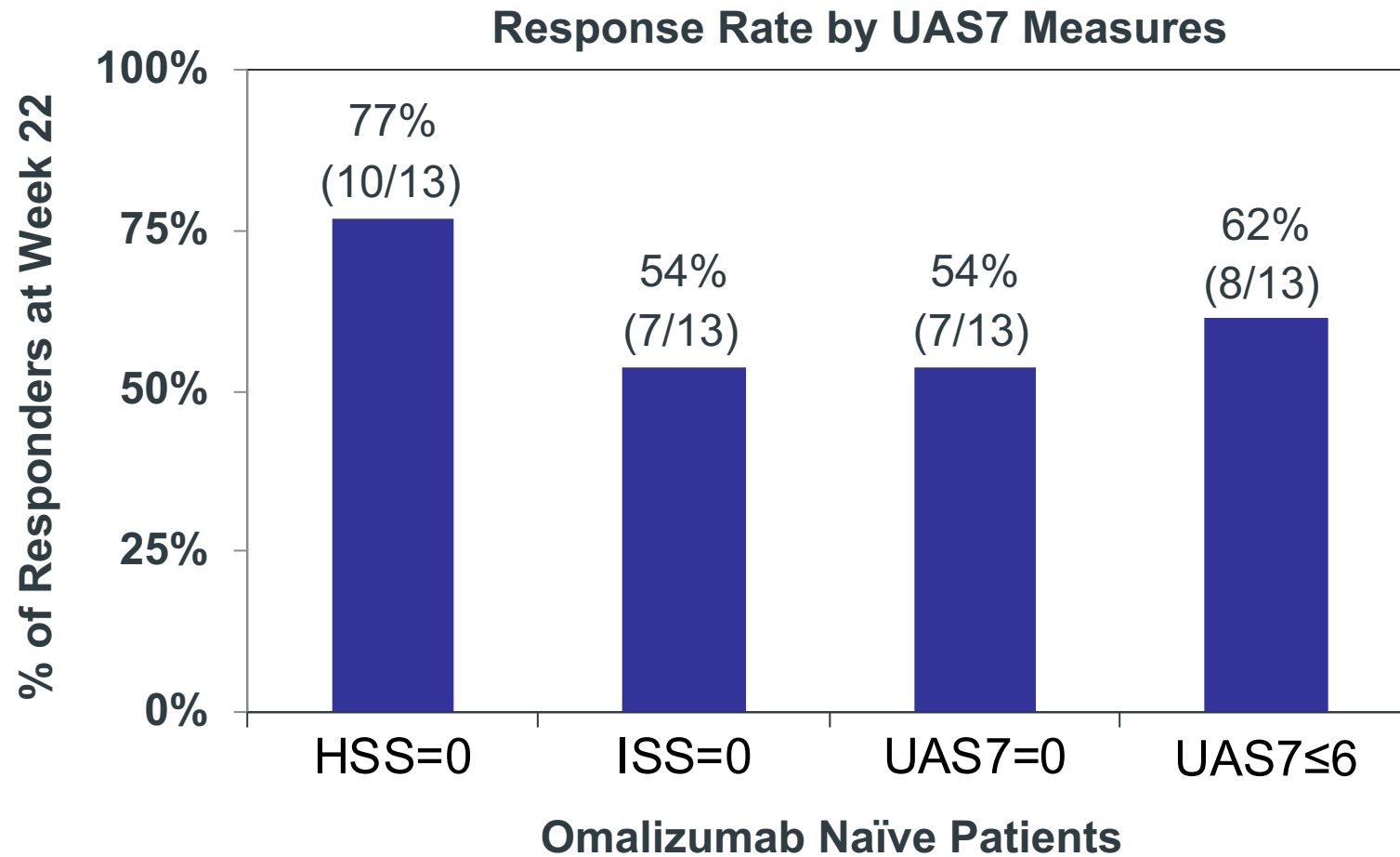
<sup>2</sup> UCT complete response:  $\geq 3$ -point improvement from baseline and score  $\geq 12$ ; partial response  $\geq 3$ -point improvement from baseline; no response  $< 3$ -point improvement from baseline



# Omalizumab Naïve CSU: Substantial Improvement in UAS7



# Omalizumab Naïve CSU: Response by UAS7 Measures



# Omalizumab Refractory CSU Cohort – Medical History

Inadequate disease control despite extensive use of omalizumab (OMA)

## Prior OMA Treatment Experience

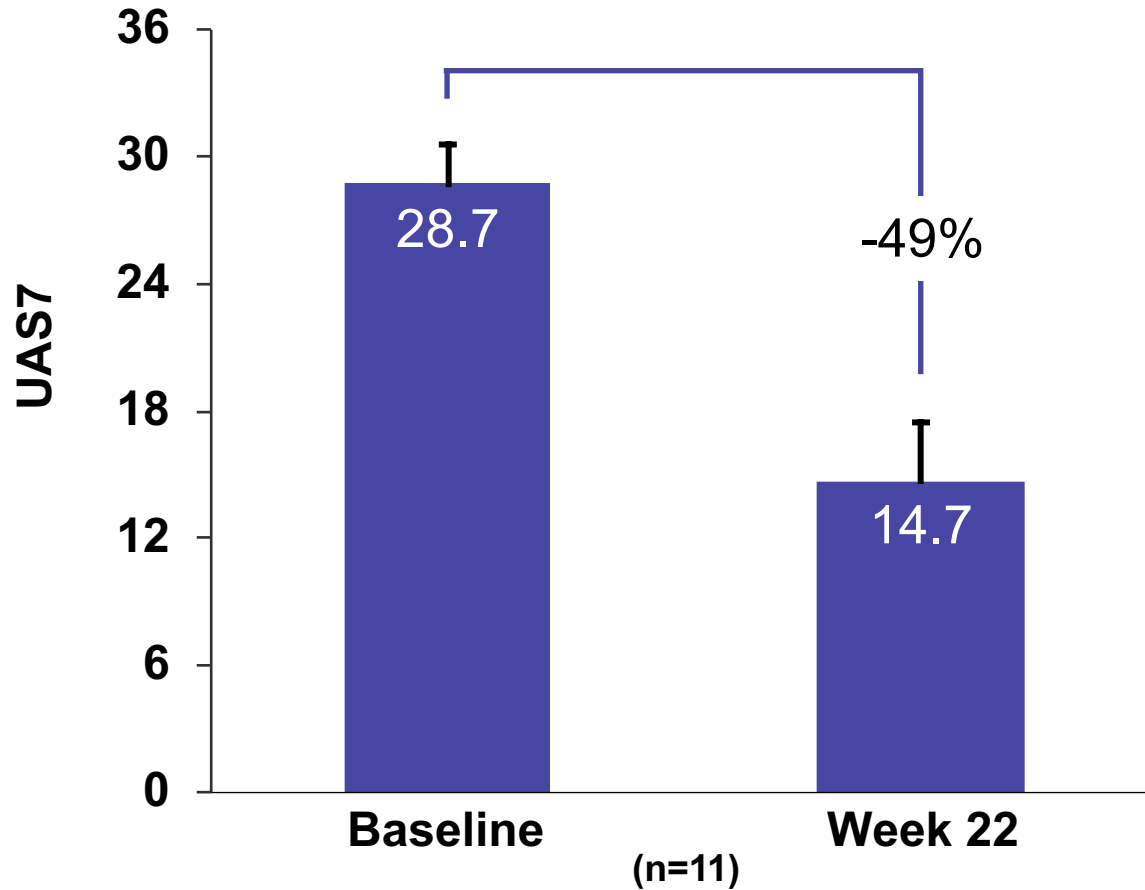
- **Average treatment duration:** ~10 months
- **Treatment regimen:**
  - Omalizumab: up to 600 mg
  - Antihistamine: up to 4x labeled dose
- **Average UCT score on Omalizumab:** 4.1

## Antolimab (AK002) Experience

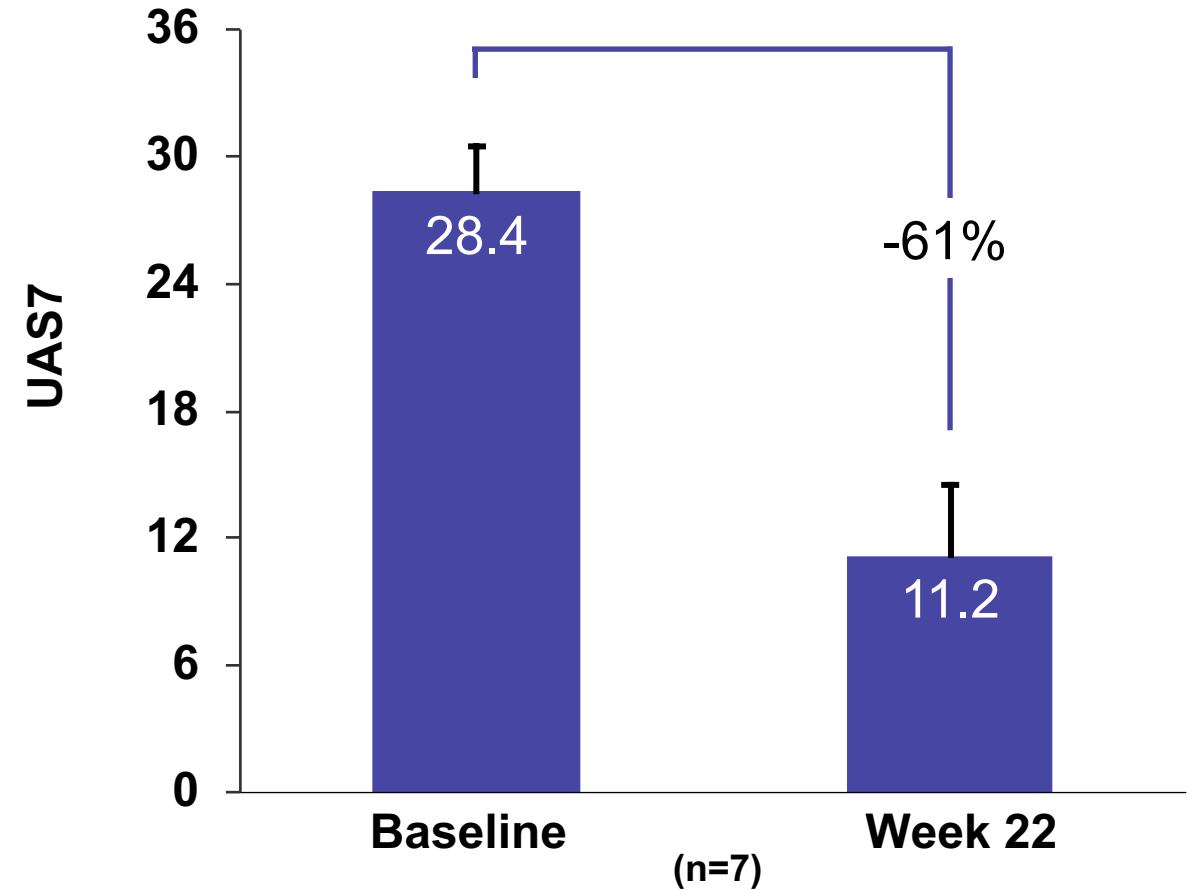
- **Time since last OMA dose:** >2 months
- **Treatment regimen:**
  - Antihistamine: up to 4x labeled dose
- **Average UCT score at baseline:** 3.7
- **Average UCT score after 6 doses:** 11.4

# Omalizumab Refractory CSU: Substantial Improvement in UAS7

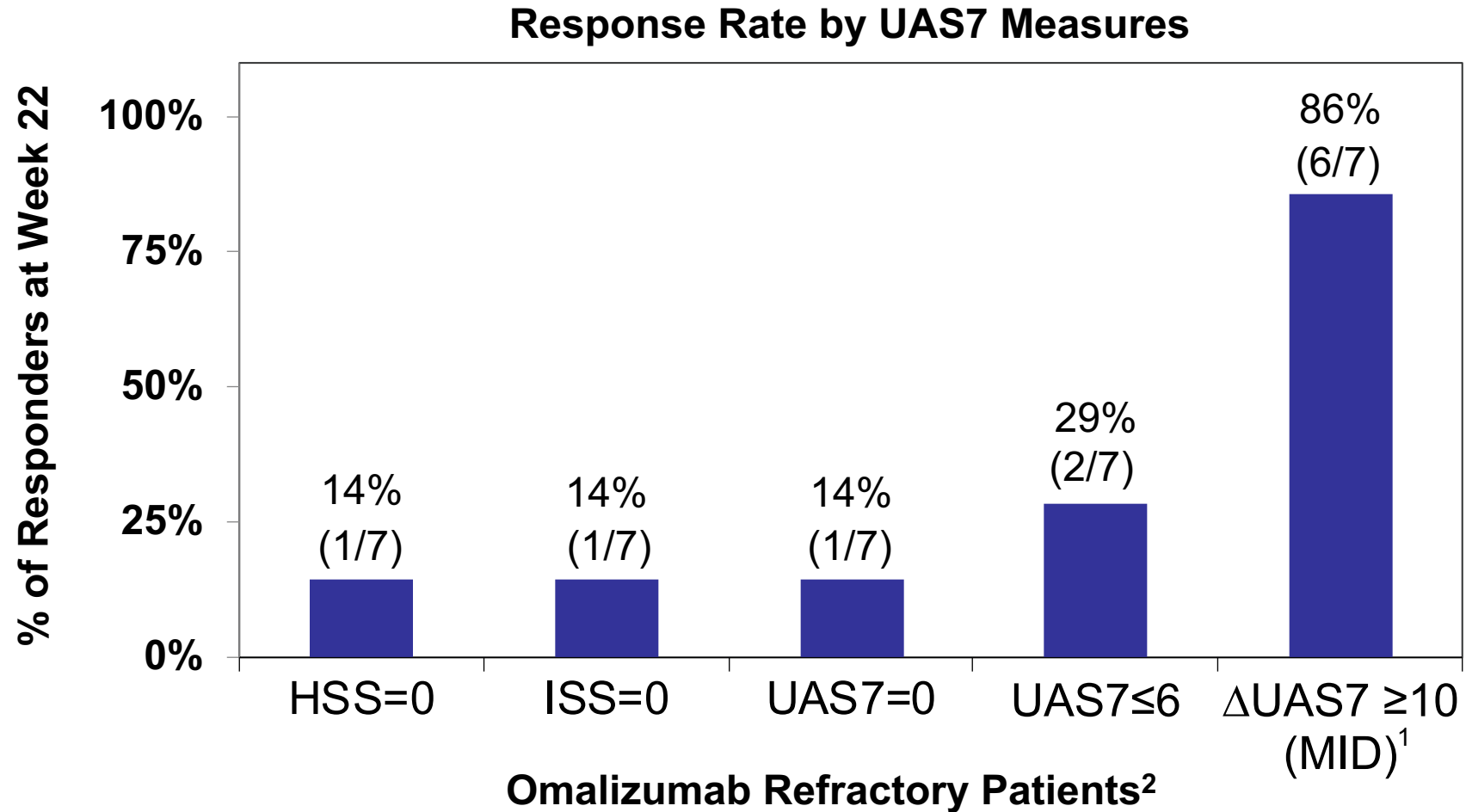
## All Omalizumab-Refractory



## Received All 6 Doses



# Omalizumab Refractory CSU: Response by UAS7 Measures



<sup>1</sup> Minimally Important Difference: Mathias, Maurer et al. Ann Allergy Asthma Immunol 108 (2012) 20-24

<sup>2</sup> Patients who received all 6 doses; In entire OMA-refractory CSU cohort, 8 of 11 patients met MID change in UAS7 ≥10

# High Response Rate by UCT with Antolimab in CIndU

## Chronic Inducible Urticaria

Urticaria Control Test Response <sup>1</sup>	Cholinergic (n=11)	Symptomatic Dermographism (n=10)
Complete Response	82%	40%
Partial Response	0%	30%
No Response	18%	30%

<sup>1</sup> UCT complete response:  $\geq 3$ -point improvement from baseline and score  $\geq 12$ ; partial response  $\geq 3$ -point improvement from baseline; no response  $< 3$ -point improvement from baseline

# Cholinergic Urticaria: 100% Response Rate by PCE<sup>1</sup> Test

	Baseline		End of Study	
	Provocation <sup>2</sup> Response < 10 mins	Number of Wheals after 30 mins	Provocation Response < 10 mins	Number of Wheals after 30 mins
CholU-1	+	21 - 50	-	0
CholU-2	+	1 - 20	-	0
CholU-3	+	1 - 20	-	0
CholU-4	+	>50	-	0
CholU-5	+	Positive	-	0
CholU-6	+	>50	-	0
CholU-7	+	>50	-	21-50

1 Pulse Control Ergometry

2 Provocation - exercise on stationary bike elevates body temperature to trigger symptoms, positive response if hives occur ≤10 minutes from start of sweating. Test terminated at 30 mins

# Safety Summary

- Generally well-tolerated
- No drug-related Serious Adverse Events
- Most common adverse event was mild to moderate infusion-related reactions (IRRs; flushing, feeling of warmth, headache, nausea, or dizziness)



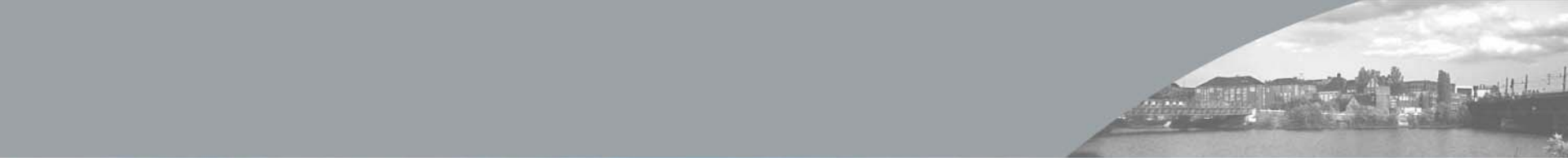
# Antolimab in Chronic Urticaria

Demonstrated activity in all forms of antihistamine-refractory  
Chronic Urticaria tested

Substantial activity in patients refractory to omalizumab

Continued activity observed in 12-month open-label extension

Antolimab is potentially a front-line biologic treatment  
for Chronic Urticaria



Thank you