



# Proof-of-Concept Phase-2a Clinical Trial of ANB020 (Anti-IL-33 Antibody) in the Treatment of Moderate-to-Severe Adult Atopic Dermatitis

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# Conflicts of interest

Advisory boards, consultancies, research grants or equity with:  
AnaptysBio, Celgene, Eli Lilly, Novartis, Janssen, Orbit Discovery, UCB Pharma

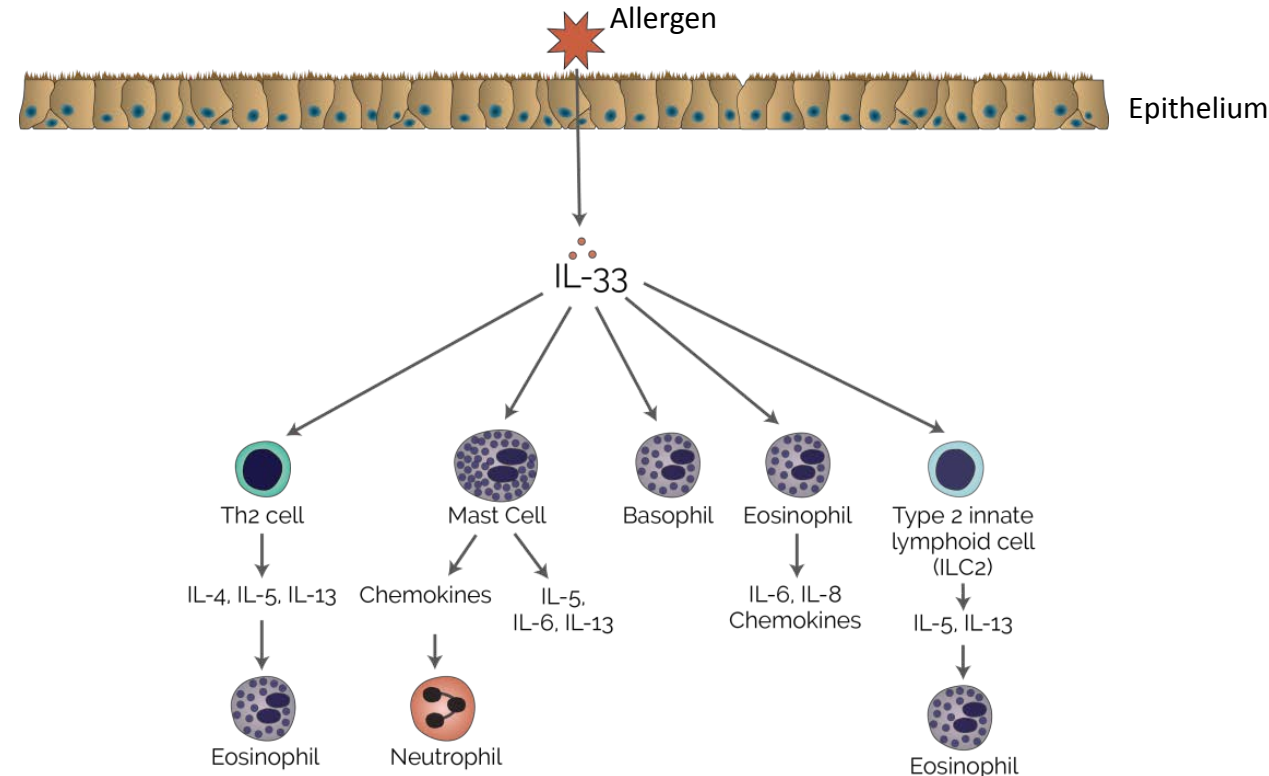
Clinical study sponsored by AnaptysBio

Travel/registration costs for AAD:  
AnaptysBio

# IL-33: Central Mediator of Type 2 Diseases

## Key Role in Pathogenesis of Atopic Dermatitis

- IL-33 is a key cytokine in type 2 inflammatory responses to allergen
  - Responsible for activation of Th2 and ILC2
  - Functions upstream of IL-4, IL-5 and IL-13
  - Modulates mast cell degranulation
- IL-33 is rapidly released by epithelium upon allergen exposure
- Genetic association of IL-33 pathway mutations with type 2 diseases<sup>1</sup>
- IL-33 is highly expressed in skin of atopic dermatitis patients with active disease<sup>2</sup>



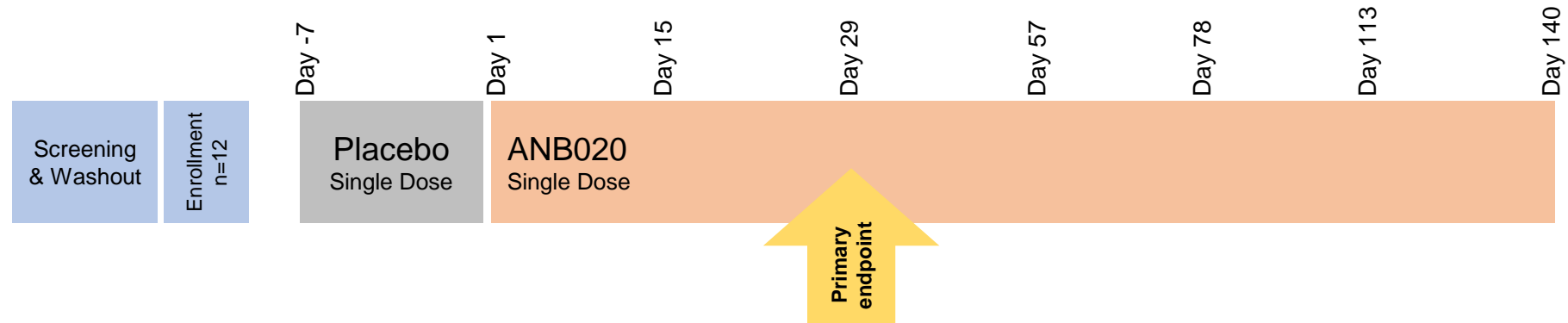
1. Ramirez-Carrozzi et al. 2014

2. Savinko et al. 2012



# ANB020 Phase 2a Atopic Dermatitis

## Proof-of-Concept Trial



- Study design:
  - Enrolled 12 moderate-to-severe adult atopic dermatitis patients inadequately controlled with topical corticosteroids
  - Single intravenous dose of placebo (Day -7) followed by a single 300 mg intravenous dose of ANB020 (Day 1)
  - EASI, 5-D pruritus, SCORAD, DLQI and IGA clinical scores determined at specific time points
- Study objective:
  - Demonstrate EASI-50 response in at least 50% of patients at Day 29 (primary endpoint)

# Baseline Characteristics

Characteristic	Average (n=12)
Age (years)	40.4 ± 13.5
Male, number (%)	11 (91.7%)
Caucasian race, number (%)	12 (100%)
Body-Mass Index	26.14 ± 4.145
EASI, score	32.25 ± 10.89
IGA, 0-5 scale	4 ± 0.74
SCORAD, score	64.79 ± 12.02
Pruritus, 5-D score	19.1 ± 4.85
DLQI, score	12.92 ± 6.54

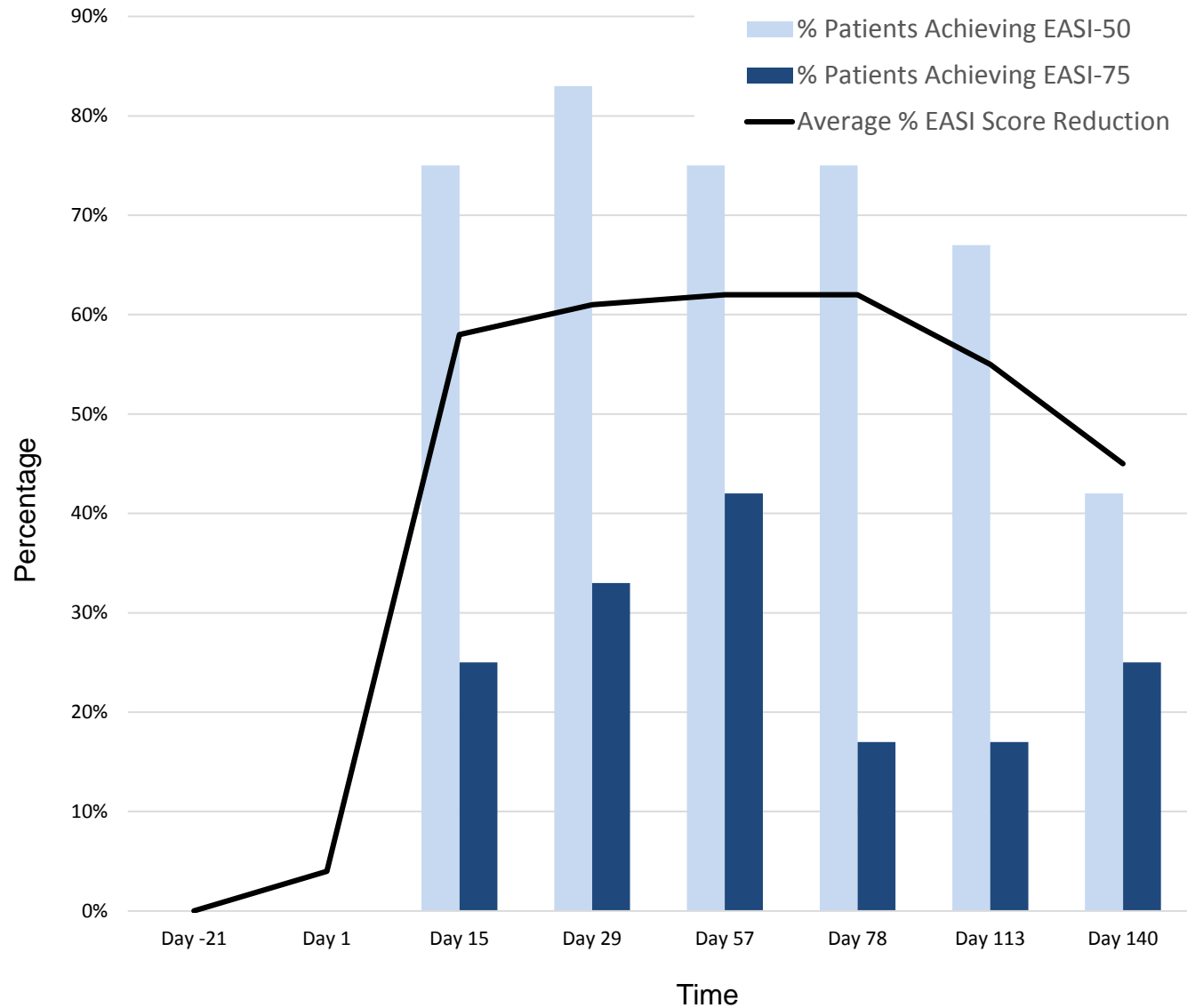
All 12 patients were inadequately controlled on corticosteroids pre-study

7 of 12 enrolled patients were treated with systemic immuno-modulators pre-study and presented with a baseline EASI score of 36

5 of 12 patients were not treated with systemic immuno-modulators pre-study and presented with a baseline EASI score of 27

# EASI Scores Following Single ANB020 Dose

Rapid response and all patients achieved EASI-50 on or before Day 57

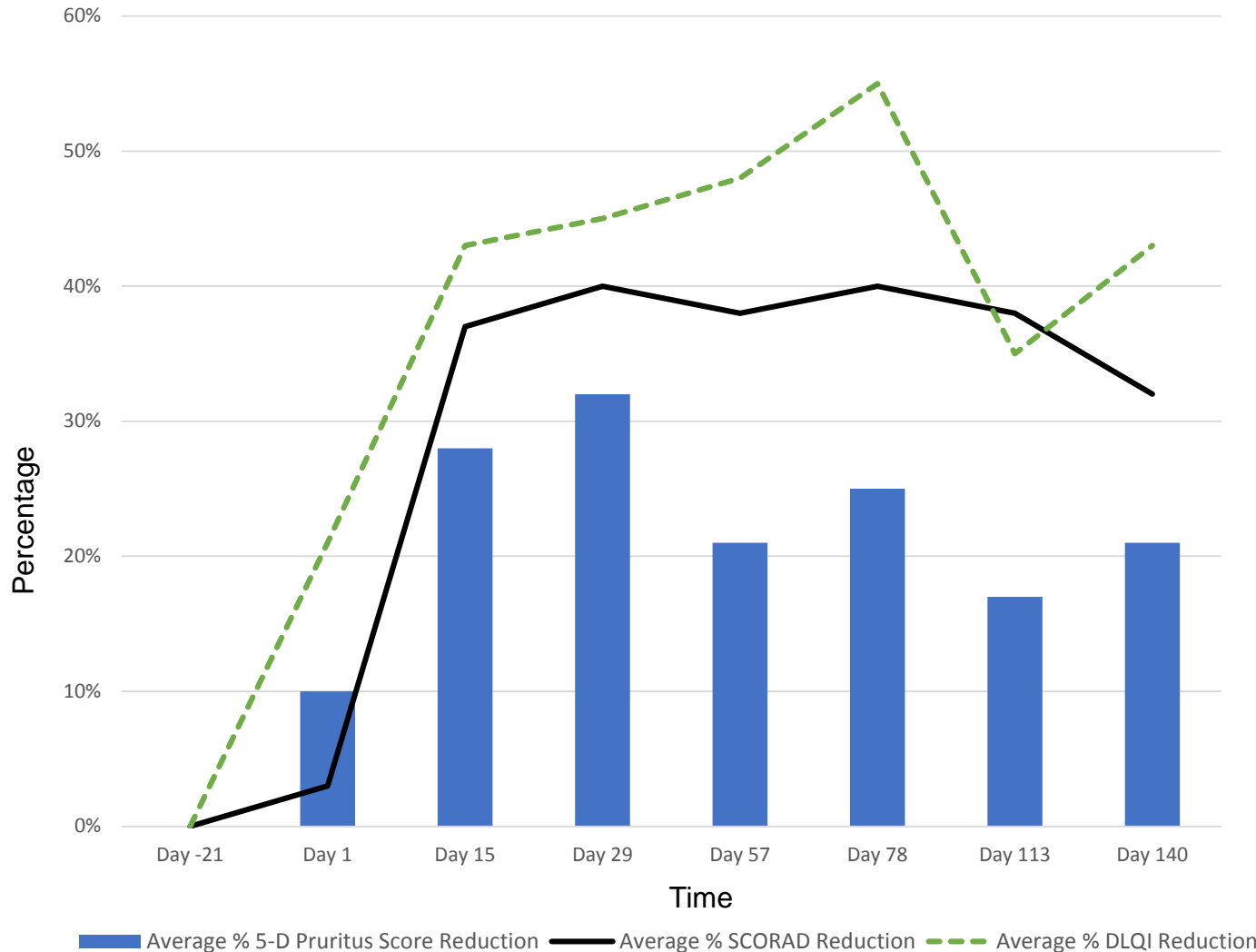


Timepoint	Average % EASI Score Reduction*	% Patients Achieving EASI-50*	% Patients Achieving EASI-75*
Day -21 (Baseline)	0%	0	0
Day 1 (ANB020 Dosing)	4%	0	0
Day 15	58%	9 of 12 (75%)	3 of 12 (25%)
Day 29	61%	10 of 12 (83%)	4 of 12 (33%)
Day 57	62%	9 of 12 (75%)	5 of 12 (42%)
Day 78	62%	9 of 12 (75%)	2 of 12 (17%)
Day 113	55%	8 of 12 (67%)	2 of 12 (17%)
Day 140	45%	5 of 12 (42%)	3 of 12 (25%)

\* Relative to baseline upon enrollment at Day -21

# Additional Efficacy Data

## 5-D Pruritus, SCORAD, DLQI and IGA Scores



Timepoint	Average % 5-D Pruritus Score Reduction*	Average % SCORAD Reduction*	Average % DLQI Reduction*
Day -21 (Baseline)	0%	0%	0%
Day 1 (ANB020 Dosing)	10%	3%	21%
Day 15	28%	37%	43%
Day 29	32%	40%	45%
Day 57	21%	38%	48%
Day 78	25%	40%	55%
Day 113	17%	38%	35%
Day 140	21%	32%	43%

\* Relative to baseline upon enrollment at Day -21

IGA scores of zero or 1 (clear/almost clear skin) observed in 25% (3/12) of patients



# Key Conclusions & Next Steps

- **Rapid and persistent efficacy following single dose of ANB020**
  - Rapid efficacy observed as early as Day 15
  - Efficacy was maximized between Day 29 and Day 57
  - All patients achieved at least EASI-50 response on or before Day 57
  - EASI responses consistent with 5-D pruritus, SCORAD, IGA and DLQI scores
- **Disease severity does not limit ANB020 efficacy**
  - ANB020 was similarly efficacious in patients with higher baseline EASI scores (treated with systemic immuno-modulators pre-study) versus lower baseline EASI score patients that did not require systemic therapy pre-study
- **ANB020 was well-tolerated and no drug-related safety signals observed**
  - Most frequent adverse event was dizziness in 17% of patients post-placebo versus headache in 25% of patients post-ANB020
  - A single serious adverse event of depression reported on Day 140 post-ANB020, which was consistent with the patient's pre-trial history of depression, and was deemed not drug-related
- **Next step: advance ANB020 into placebo-controlled, double-blind, randomized 200-300 adult moderate-to-severe atopic dermatitis Phase 2b trial**
  - Assess different dose levels and dosing frequencies of subcutaneously-administered ANB020



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