

Corporate Overview

NASDAQ: ORKA

March 2025



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Fully funded through a potential psoriasis breakthrough

2024 was a year of superb execution...

Better biologics overdeliver in PsO

...and continued external tailwinds

-
- UCB's Bimzelx launch exceeding expectations \$1.4B¹ 2025 and \$5B+ peak sales consensus

Raised >\$475M in two oversubscribed transactions and went public via reverse merger

Lead program ORKA-001 entered the clinic in December 2024, well ahead of schedule

- Skyrizi continues to exceed forecasts now projecting \$12.5B 2027 sales in psoriatic disease
- Orals do not reach biologic efficacy e.g., JNJ-2113 (icotrokinra) Ph3 in Q4 2024

Clinical data catalysts coming every 6 months

- 2H 2025 ORKA-001 HV PK
- 1H 2026 ORKA-002 HV PK
- 2H 2026 ORKA-001 PsO Ph2a
- Beyond ORKA-002 Ph2, ORKA-021: major additional sources of optionality and upside

Multiple "ways to win"

- Ultra-long dose interval (1-2x per year)
- Highest IL-23 PASI 100 (à la KNOCKOUT)
- Long-term "remissions" in some patients

Fully-funded through 2027

- Funded well-beyond ORKA-001 PsO Ph2a readout, with cash through 2027 no need to raise
 on healthy volunteer data
- 55.1M total shares of common stock and common stock equivalents



On a mission to enable freedom from chronic skin disease

Our goal

Help patients with chronic skin conditions experience the greatest possible freedom from disease



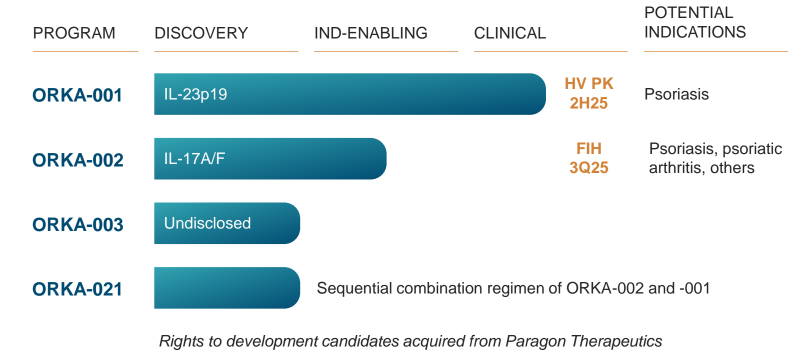
Highest possible rates of disease clearance



Fewest number of doses

Our approach

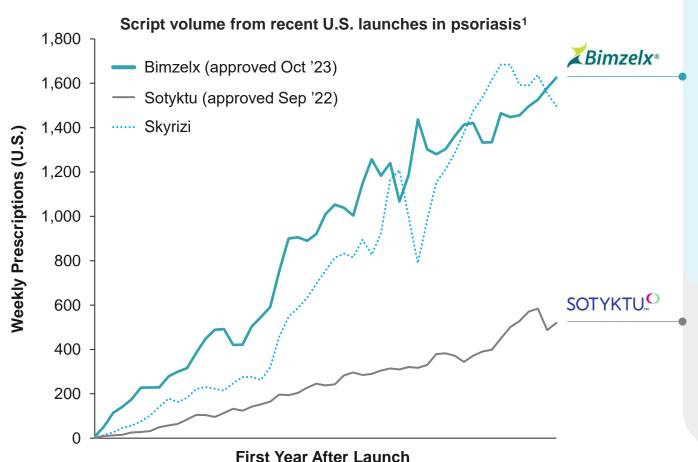
Advance potentially **best-in-class antibodies** targeting mechanisms with **proven efficacy and safety**





Bimzelx launch shows that better biologics will win in psoriasis

Bimzelx versus Sotyktu performance validates our thesis



- UCB's Bimzelx launch has exceeded expectations, driven by strong demand – consensus sales of \$1.4B² in Year 2 and \$5B+ peak
- Market underestimated the opportunity UCB market cap \$15B pre-launch vs. \$35B one year later (\$20B market cap created on Bimzelx alone)
- Strong launch driven by PsO in U.S. proof point that smaller, non-incumbent company can effectively commercialize in PsO
- Sotyktu underperformed due to lack of demand sub-optimal efficacy with JAK-like safety overhang
- Market access dynamics not meaningfully different from Bimzelx – not a major driver



The psoriasis market will continue to reward biologic innovation

Psoriasis is a massive, growing market where biologics are still underpenetrated

\$30B+ growing market today with further potential in mild-to-moderate disease

Pharma believes in the future of this market

Shown by major investments in orals, which sacrifice efficacy in favor of perceived convenience













Dermatologists and patients want new and better biologics

Even a base-case profile of Q6M dosing with equal efficacy surveys at ~50% of new patient starts in 2030+

Differentiated biologics have a proven ability to gain market access

Bimzelx forecasted to reach peak sales of \$5B+



Showing smaller, non-incumbent companies can achieve access



ORKA-001 & -002 complement each other to address all PsO/PsA

ORKA-001

For patients with purely skin disease



Majority of dermatologists prefer an anti-IL-23p19

ORKA-002

For patients with joint involvement, including PsA, or recalcitrant skin disease



Anti-IL-17 preferred, and IL-17A/F emerging as the best approach

ORKA-021

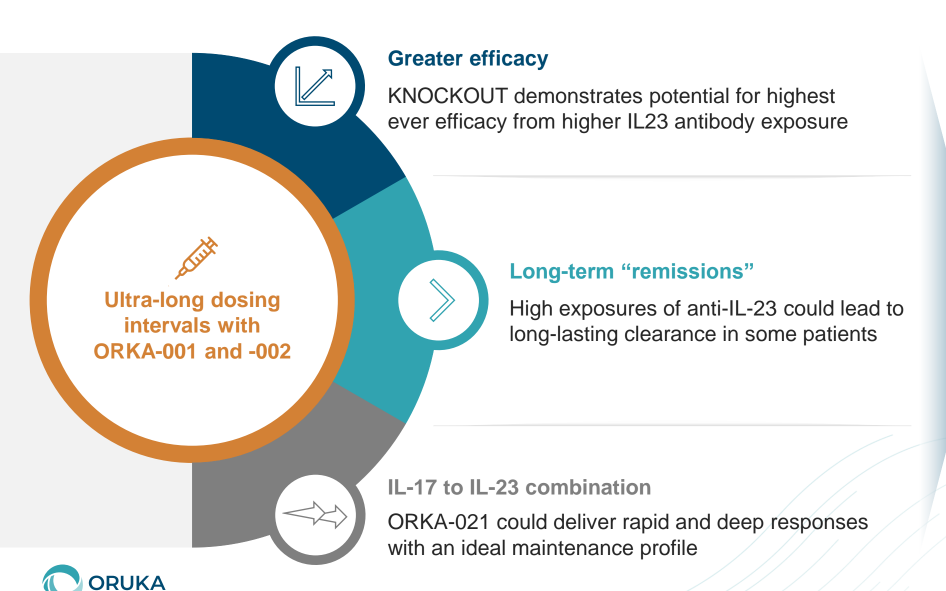
Sequential **combination of -002 and -001 –** rapid response with ideal maintenance profile



Creates another way to "win" in defining the best possible regimen in PsO and PsA



Multiple orthogonal paths for Oruka to maximize differentiation



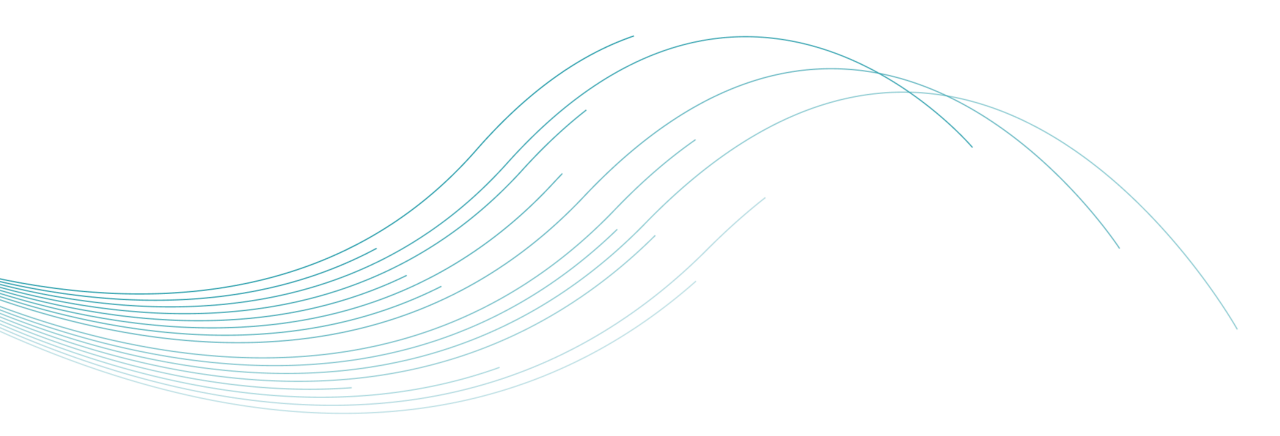
Maximizes odds of having a strong value proposition to achieve preferred access and price for innovation

Clinical data catalysts coming every 6 months going forward

	2025		2026		Beyond
ORKA-001	FIH Ph1 ☑	Interim PK in HVs	Final PK in HVs	Ph2a in PsO: PASI 100 & response duration	Major clinical
ORKA-002		FIH Ph1 (3Q25)	Interim PK in HVs	Final PK in HVs	catalyst planned every six months

Fully funded through 2027, >1 year past ORKA-001 Ph2a readout in PsO

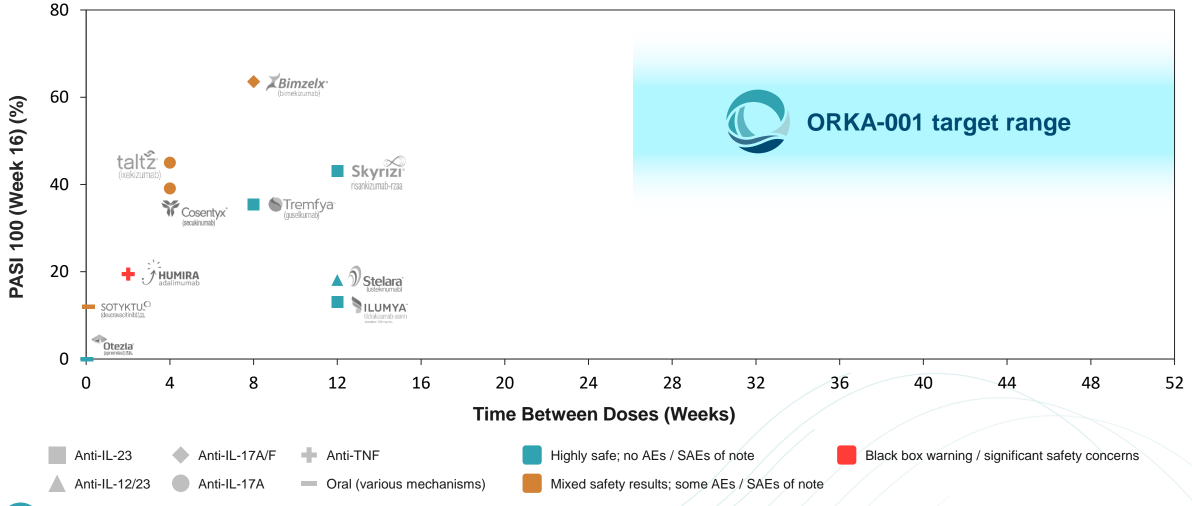




ORKA-001: potentially best-in-class anti-IL-23p19



Biologics have raised the bar on standard of care in PsO, but there is ample room for improvement





Base case is best-in-class, upside could be paradigm changing

Dosing interval

Efficacy

Base case scenario

Once per six months

Comparable PASI 100 to Skyrizi

Best-in-class profile

Upside scenario

Once per year and/or patient-specific

Better PASI 100 than Skyrizi

Paradigm-changing



ORKA-001 could be the last word in IL-23p19 inhibitors

Binds similar epitope to Skyrizi (risankizumab) with similar potency

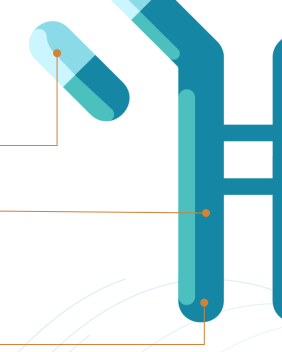
- Validated mechanism of action
- Specific for IL-23p19 (not IL-12/23 p40)
- $K_D < 5 pM$
- Predicted equivalent safety
- Predicted to meet or beat efficacy

Novel IP for composition of matter into 2040s

Half-life extension through validated Fc modification (YTE mutations)

- Higher exposure to increase efficacy
- Longer exposure to reduce dosing frequency

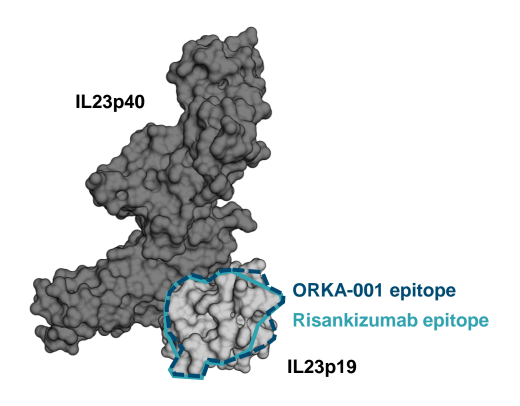
Effector-null human IgG1 Fc





ORKA-001 binds a similar epitope as risa with similar potency

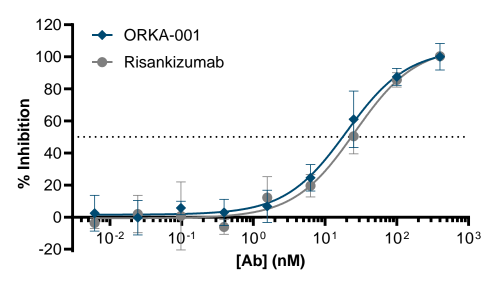
ORKA-001 binds a nearly identical epitope to risankizumab



Comparable affinity (<5 pM) as well

ORKA-001 shows comparable potency to risankizumab

Inhibition of IL-17 release in human PBMCs



	ORKA-001	Risankizumab	Guselkumab
Relative IC50	0.76	1.0	4.83

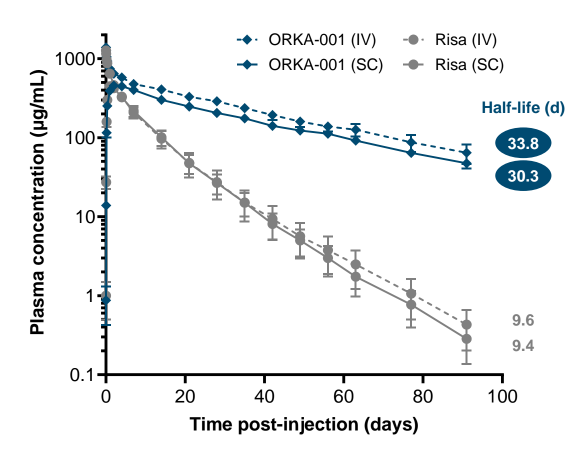
Similar results observed across a range of in vitro assays

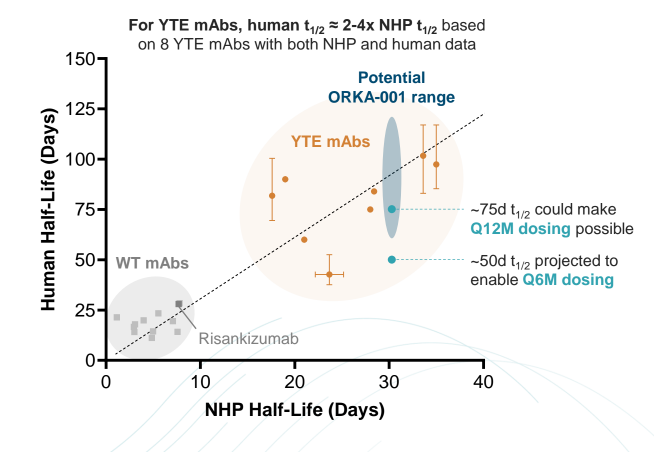


Clinical experience with YTE predicts significant t_{1/2} extension

ORKA-001 has a >3x longer half-life than risankizumab in NHPs

Implies ORKA-001 could have a significantly longer half-life in humans



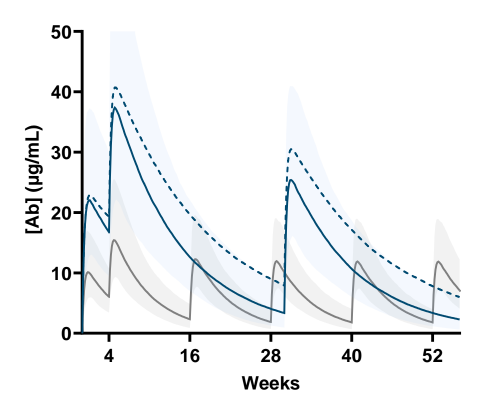


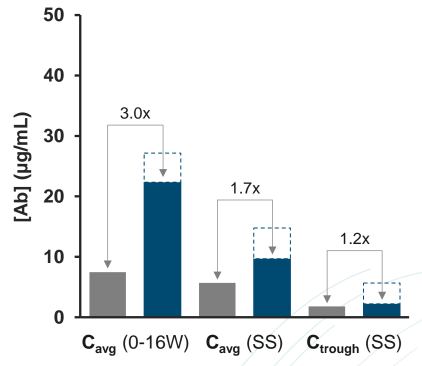


Base case is achievable even at lower end of predicted half-life

ORKA-001 exposure could exceed Skyrizi under a variety of half-life scenarios

- **ORKA-001 (~50d half-life):** 300 mg W0, 4, Q6M **Skyrizi:** 150 mg W0, 4, Q12W (approved regimen)
- --- ORKA-001 (~75d half-life): 300 mg W0, 4, Q6M





Even at a 50-day half-life, Q6M dosing with ORKA-001 is projected to give a significantly higher C_{avg} and C_{trough} than Skyrizi



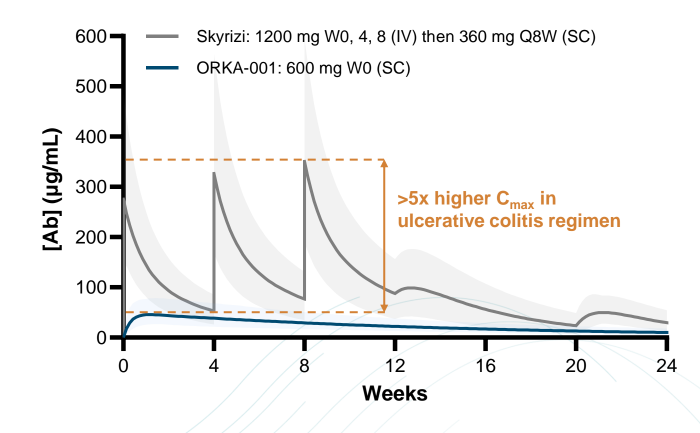
ORKA-001 benefits from a large body of clinical evidence with IL-23 inhibition

Very uncommon to have clinical precedent in large numbers of patients for the safety of higher exposures

- Peak and average exposures of ORKA-001 dosed at 600 mg are multiples lower than those with approved Skyrizi regimens in IBD
- No correlations observed at the patient level between exposure and safety signals across
 >4,000 patients dosed with Skyrizi in clinical trials

All five IL-23p19 inhibitors with published data in psoriasis have performed as expected based on their biophysical properties

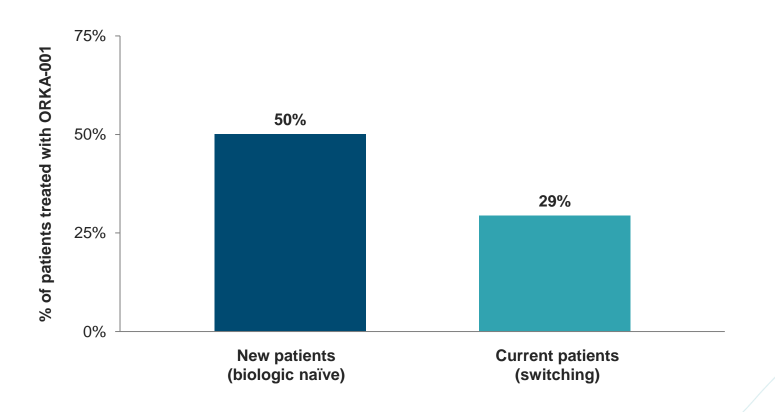
Skyrizi regimen in UC establishes the safety of very high exposures





Dermatologists view the "base case" as highly attractive

In the "base case," dermatologists would put half of new patients on ORKA-001 even when accounting for entry of new oral medicines



Multiple examples support dosing as a major commercial differentiator:

PsO ►Tremfva[®] VS. 'auselkumab)' Q12W Q8W **Fasenra** Nucala 🌶 **Asthma** VS. (mepolizumab) Q8W Q4W **EYLEA wAMD** VS. Q8W Q4W

Increasing excitement about drugs with long dosing intervals:

- Positive Phase 3 results for depemokimab (GSK), ocrelizumab (Roche), lenacapavir (Gilead), all given twice-yearly
- GSK acquired Aiolos for a long-acting YTE mAb targeting TSLP



Three potential upside scenarios for ORKA-001





Higher exposure could drive higher PASI 100



1-year dosing interval

Enabled by half-life extension



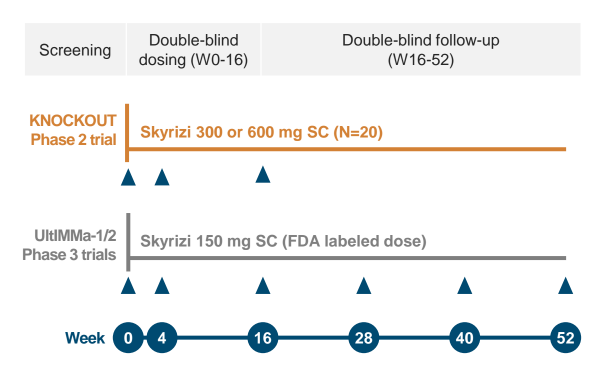
Disease modification

Patient-specific dosing to allow for treatment-free remissions

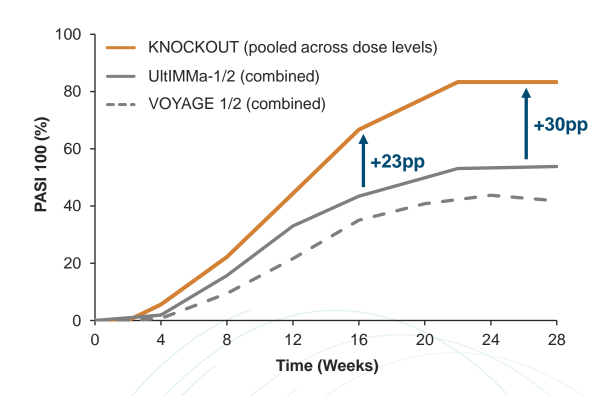


Higher exposures drove higher efficacy in KNOCKOUT study

KNOCKOUT evaluated 2-4x the approved Skyrizi dose...



...and resulted in the highest PASI 100 rates observed to date

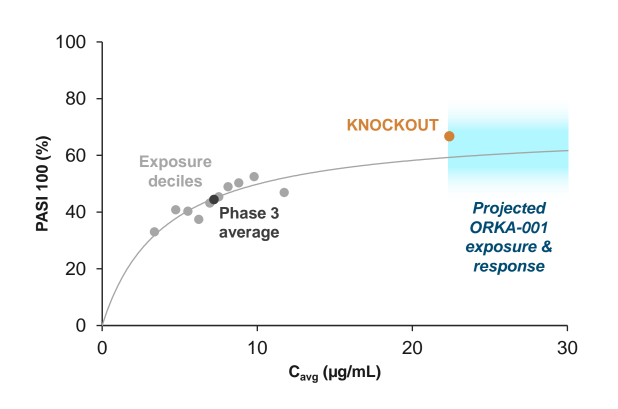


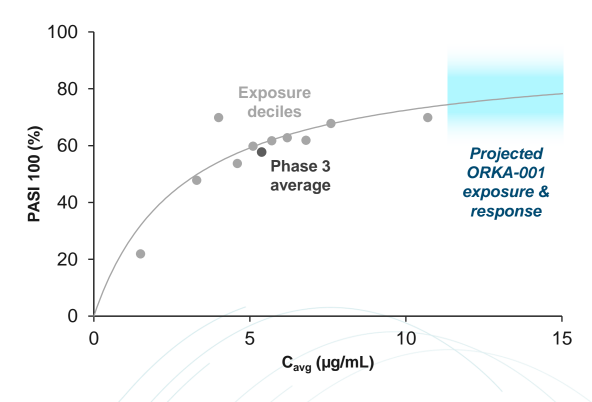


ORKA-001 could drive higher efficacy based on KNOCKOUT and a consistent exposure-response trend across trials

Risankizumab induction phase (0-16 weeks)

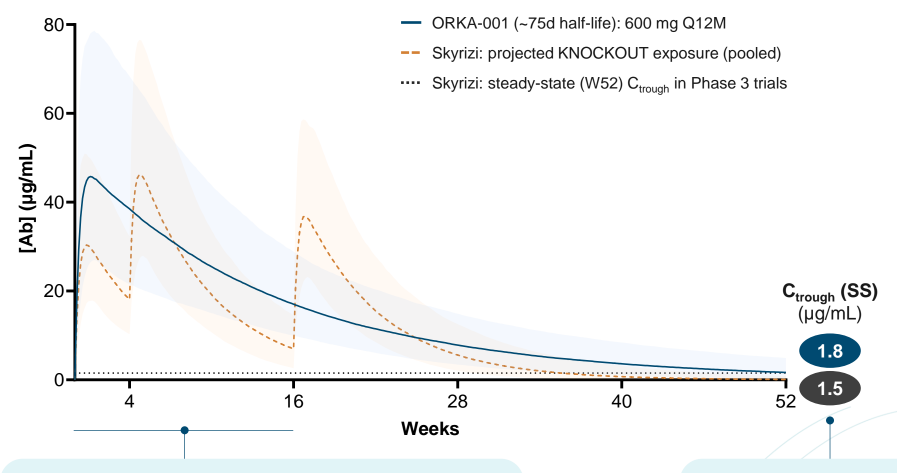
Risankizumab steady-state phase (40-52 weeks)







KNOCKOUT-like exposures are possible with one dose per year



With a ~75-day half-life, ORKA-001 at one dose per year could match or exceed both KNOCKOUT early exposures and steady-state trough levels of standard Skyrizi dosing

Comparable or greater early exposure vs. KNOCKOUT

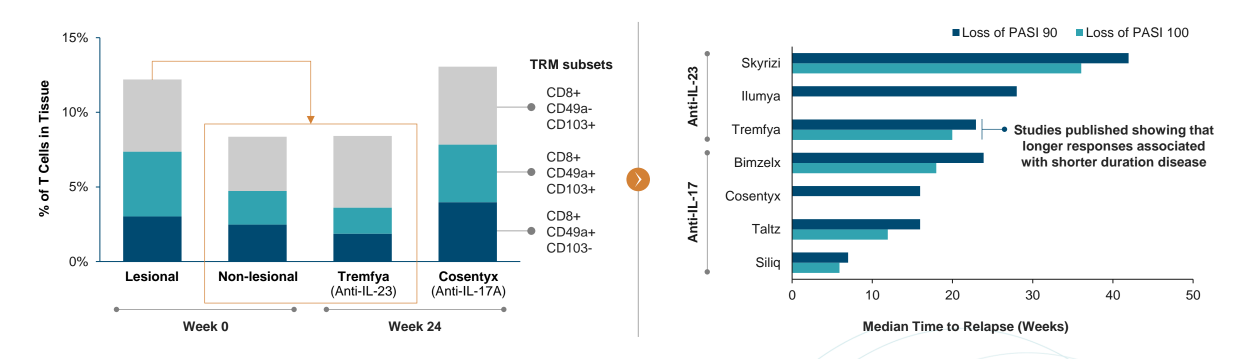
Comparable or greater steady-state C_{trough} vs. Skyrizi



Potential for disease modification or cure by depleting TRMs

Anti-IL-23 acts upstream of disease-causing TRMs, and has a unique ability to deplete them from tissue

Leading to long-lasting remissions after treatment withdrawal – pointing to possibility of disease modification



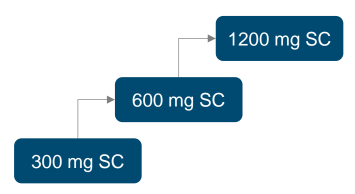
Excitement growing in dermatology community to **test disease modification potential of high anti-IL-23** exposures early in disease — **a perfect opportunity for ORKA-001**



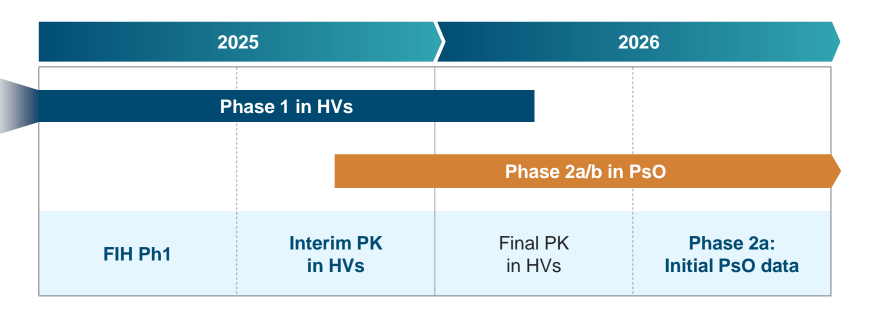
ORKA-001 development path sets up a catalyst-rich next 2 years

Phase 1 study to evaluate the safety, tolerability, and PK of ORKA-001 in healthy participants

 Placebo-controlled, single ascending dose study (NCT06698939)



- Conducted at a single center in New Zealand
- ~24 healthy volunteers



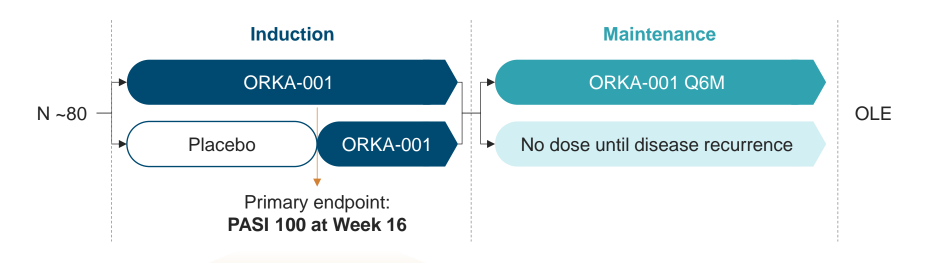
Potential for rapid de-risking, value recognition, and path to BLA

- Interim PK is highly validating, showing both basis for differentiation and early safety
- Validated clinical endpoints show highly robust correlation between Phase 2 and 3
- Rapid timelines possible in PsO average time from FIH to BLA/NDA is 6.5 years



Phase 2a to test KNOCKOUT-like exposures of ORKA-001

Phase 2a proof-of-concept study of ORKA-001 in moderate-to-severe psoriasis

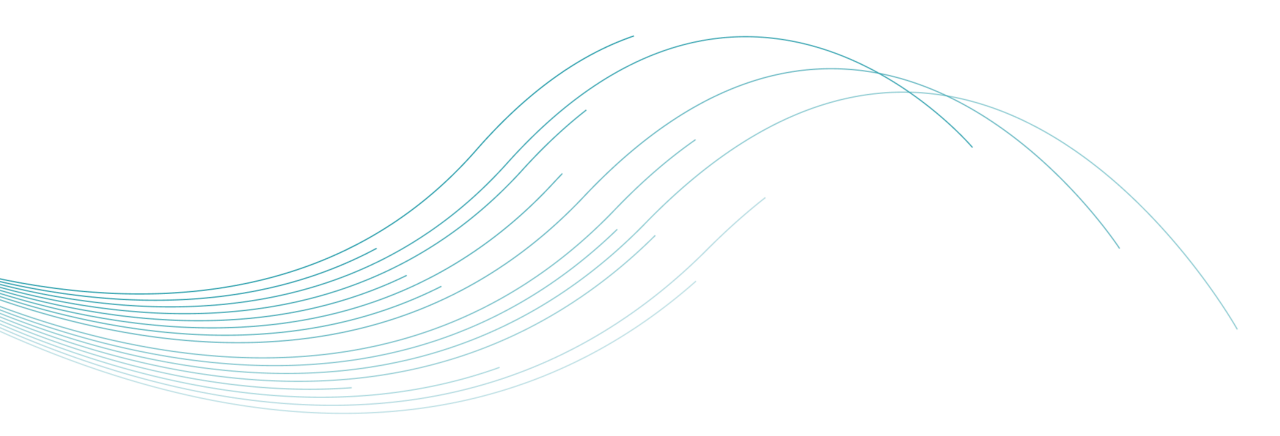


- Phase 2a expected to start in 2H 2025
- Initial efficacy data expected in 2H 2026
- Phase 2a/b trial enables efficient transition to Phase 2b dose-ranging study following Phase 2a

Initial data readout has potential to inform all "upside" scenarios:

- Efficacy at W16 for all patients (as well as later timepoints for some patients)
- Preliminary durability, indicating the potential for extended dosing intervals and longer-term remissions



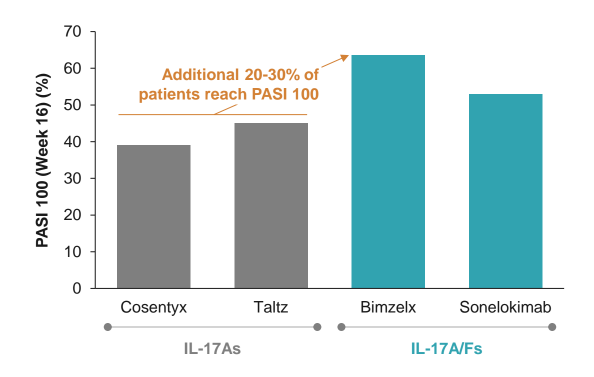


ORKA-002: potentially best-in-class anti-IL-17A/F



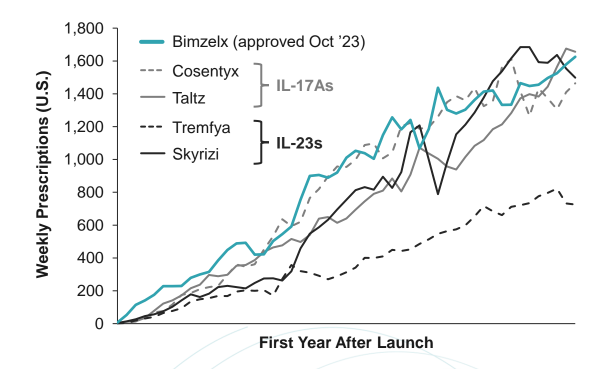
IL-17A/F dual blockade has emerged as the superior strategy

IL-17A/F shows superior efficacy to IL-17A in PsO



Superior efficacy in other indications as well

Bimzelx has had a very strong launch, validating both IL-17A/F and the ability to differentiate in PsO

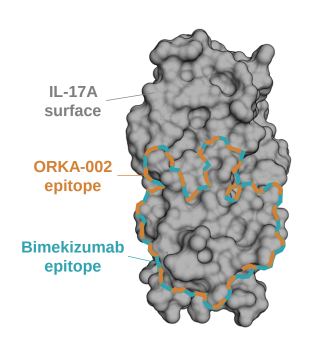


Bimzelx consensus peak sales estimate of >\$5B

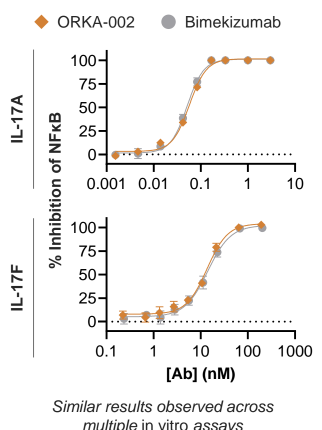


ORKA-002 has a dramatically extended half-life vs. bimekizumab

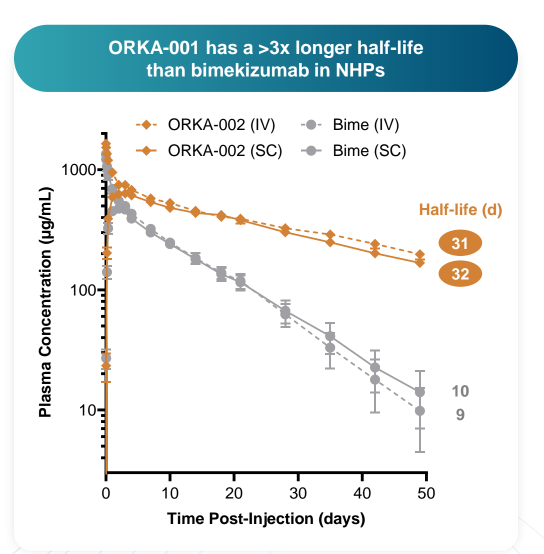
ORKA-002 binds a nearly identical epitope to bimekizumab with comparable potency



Nearly identical epitope for IL-17F as well Comparable picomolar affinity for IL-17A and IL-17F



multiple in vitro assays

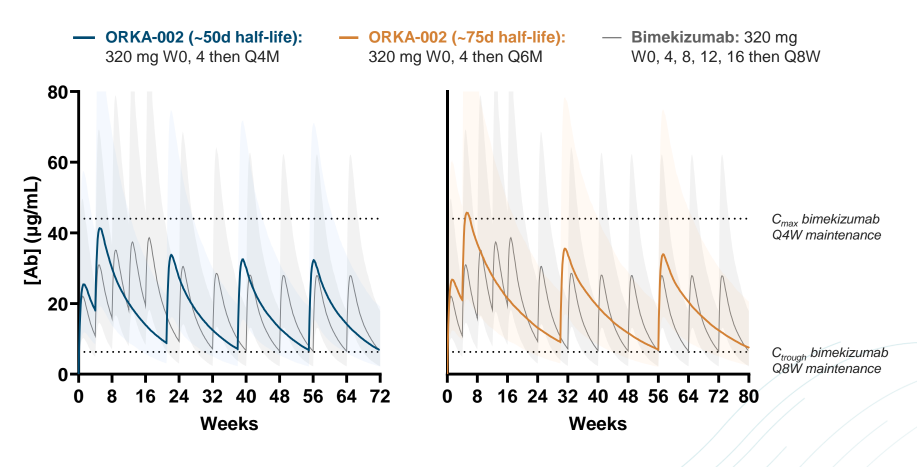




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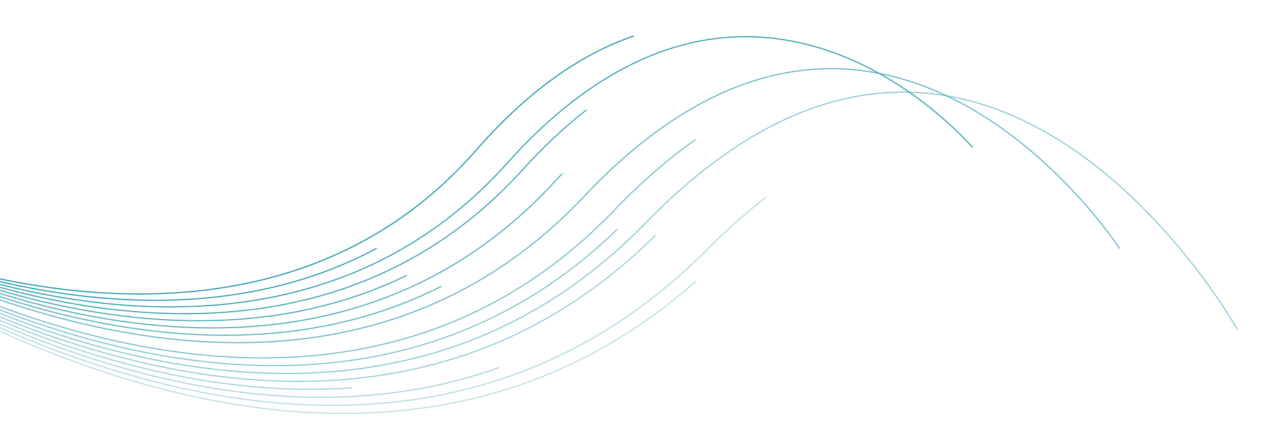
Potential for 2-3 doses per year enabled by half-life extension

Projected C_{trough} of illustrative ORKA-002 regimens exceeds approved bimekizumab regimen



A ~50-day half-life could enable Q4M dosing and ~75-day half-life could enable Q6M dosing while maintaining trough antibody levels above bimekizumab





ORKA-021



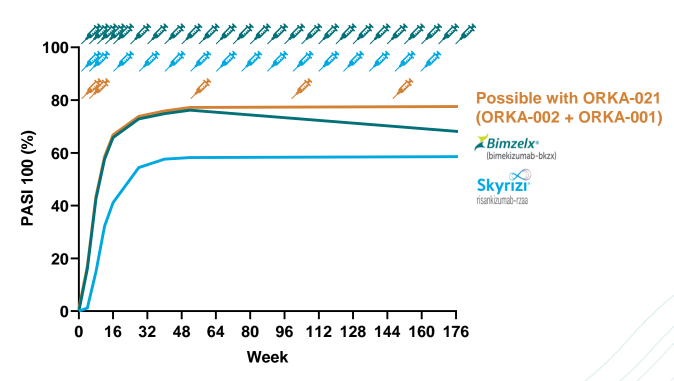
ORKA-021: Potential to combine the best of IL-17s and IL-23s

IL-17s: fastest onset and highest peak response



IL-23s: less frequent dosing and best durability and safety

Combining the two mechanisms sequentially could provide the "best of both worlds"



Feedback from U.S. dermatologists:

"It really sounds like a great option"

"Conceptually beautiful"

"The only reason this hasn't been done is that no company has both"





Shares outstanding

s of Feb 15, 2025		Number of shares ¹
Common stock	Shares outstanding	37.4M
Common stock	Preferred stock (as-converted to common stock)	11.4M
equivalents	 Pre-funded warrants 	6.2M
Common stock and		
common stock	Total outstanding ²	55.1M
equivalents		

