

Corporate Presentation

June 2025

Rĕclūdo (Latin)

transitive verb III conjugation

1 to open

2 to open up, to disclose, to reveal, to unlock



Unlocking New Therapeutic Possibilities

Unlocking Previously Undruggable SH2 Domains of High Value Targets in Inflammatory Diseases



Differentiated candidates in validated pathways & blockbuster markets



STAT6 inhibitor

- Sanofi partnership
- Validated by IL-4Rα and IL-13 biologics



BTK SH2 inhibitor

- Wholly-owned
- Validated by BTK kinase inhibitors

Undisclosed Programs

 Additional SH2 domain high value targets

OPERATIONAL STRENGTH

Experienced drug developers | Well-capitalized to value-driving milestones Sanofi option: 50-50 US profit share

PROPRIETARY PLATFORM DRIVES OPTIONALITY

120 human SH2 domains; coveted targets previously considered "undruggable" Fuels robust pipeline and multiple partnering opportunities

Experienced Leadership Team





Nancy Whiting, Pharm.D. President and CEO



Ajay Nirula, M.D., Ph.D. EVP, Head R&D



Matt Caldemeyer, **MBA CBO**



Catherine Bovenizer, C.P.A SVP, Finance



Adcetris®, Tukysa®, Padcev®, Tivdak®















Rituxan®, Tecfidera®, Silig®, Taltz®, Olumiant®, Omvoh®, Ebglyss®









GeneFormatics



Brian Hodous, Ph.D. SVP, Chemistry



Daniel Treiber, Ph.D. SVP, Discovery **Technology**



Paul Smith, Ph.D. SVP, Biology



Nick Lydon, Ph.D. Co-founder. **Board Member**

























Kinetix Novartis/CIBA-GEIGY

Gleevec® Lasker-DeBakey Award, Japan Prize

Ayvakit™

Vanflvta[®]

OpzeluraTM, Gilenya [®], Kesimpta[®]

\$102M Series A







Nick Lydon

Strategic Collaboration with Sanofi for STAT6 Inhibitor



50%

Option to participate in 50% U.S. profit share

>\$1.3B

Upfront payments and potential future development and sales milestones

Significant royalties

Up to double-digit royalties on future sales

Recludix

- Preclinical research and early clinical development
- Certain U.S. co-promotion activities

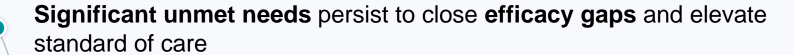
sanofi

- From Phase 2 onwards, WW clinical development
- U.S. co-promotion and ROW commercialization

Significant Unmet Medical Needs in Inflammatory Disease Remain



>60 M patients diagnosed globally each year with immune-related inflammatory disease; potential for rapid market expansion



Risk of **infection** and **other serious events** with current therapies pose major **safety barriers** that limit addressable population

Self-injection can be **burdensome** and **reduces compliance for many patients**

A diverse range of sub-populations have **no suitable therapeutic option** and remain untreated

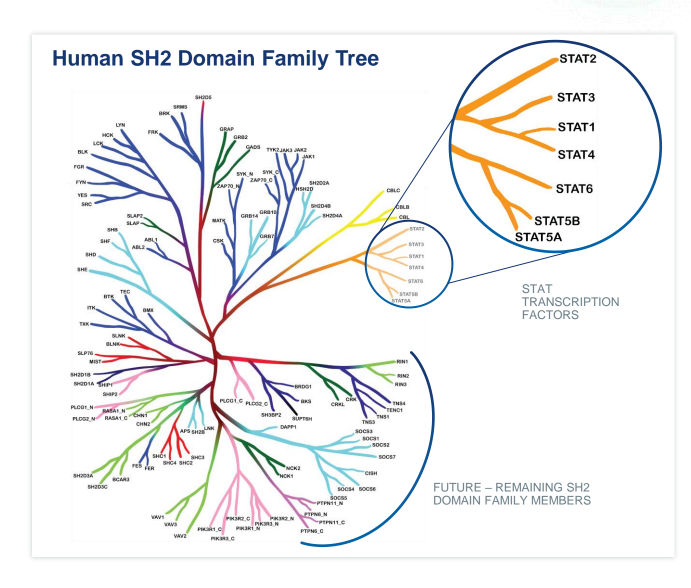
SH2 Domains Have Previously Been Deemed "Undruggable"



Significant opportunity in targeting SH2 domain proteins

Src Homology 2 (SH2) domains are highly conserved protein domains that have long been recognized as attractive drug targets

- Small protein modules made up of ~100 amino acids
- 120 human SH2 domains
- Play a key role in mediating protein-protein interactions
- The SH2 domain of STAT proteins is required for:
 - Binding to cytokine receptors
 - Dimerization of STAT proteins



Molecular Cell (2006) vol. 22, p.851

Recludix Platform: Integrated Proprietary Technologies & New Chemical Approaches





Custom SH2 Domain Targeting DNA-Encoded Libraries

Discovery and SAR DELs

Hundreds of millions of molecules

Proprietary DEL Selection & SAR Analysis

Rapid massively parallel determination of structure-activity relationships



Prodrug Modality Expertise

Drives enhanced target coverage

High Quality Lead Compounds

Structure-Based Design

Proprietary co-crystallography systems



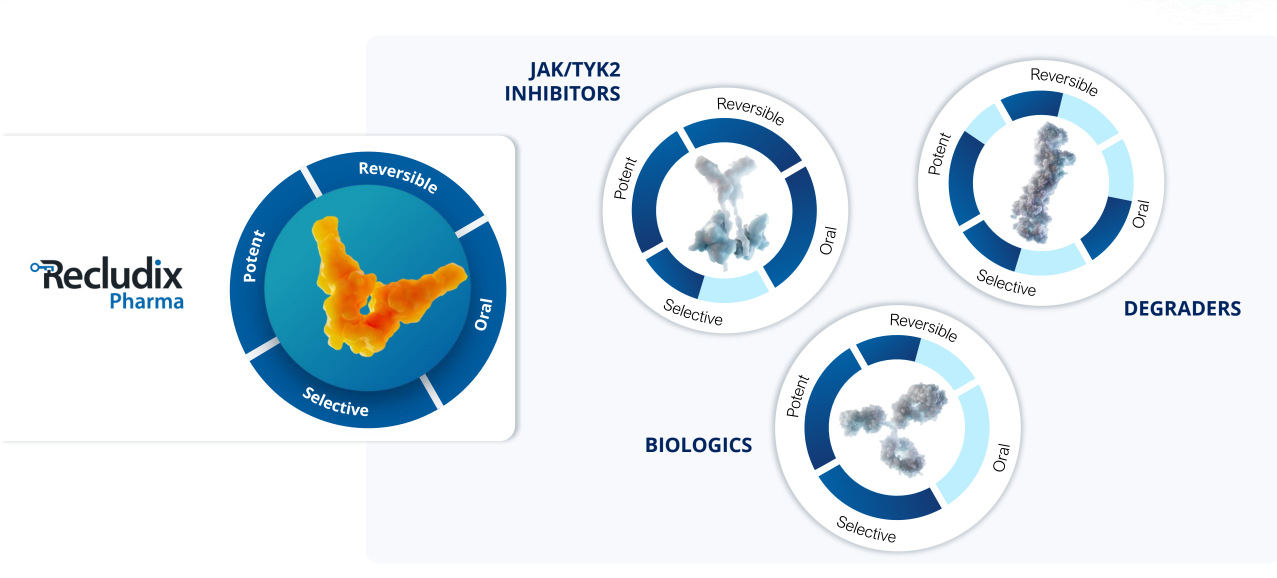


SH2 Domain Screening Tool

Unique panel of Drives SH2 domain assays selectivity

STAT SH2 Domain Inhibition Enables a Best-in-Class Product Profile



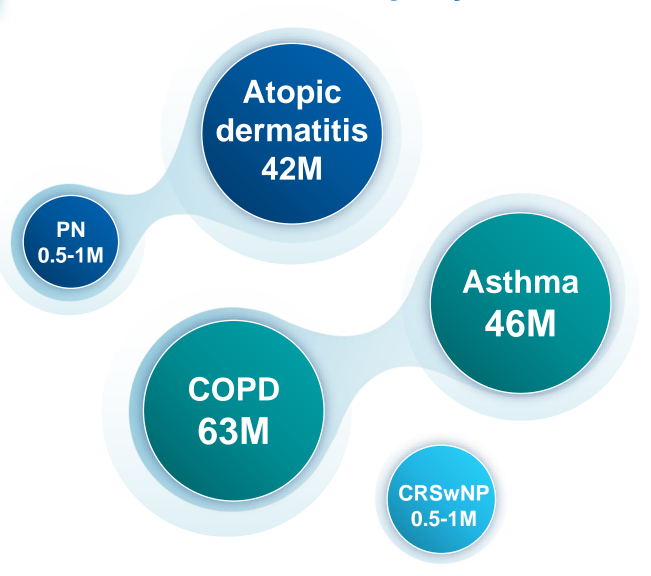




STAT6

Opportunity to Serve Large Patient Populations in High-Value Markets with a Uniquely Differentiated Product





>150M TOTAL PATIENT IMPACT

LARGE POPULATIONS

Targeting broad, established patient populations with persistent unmet needs

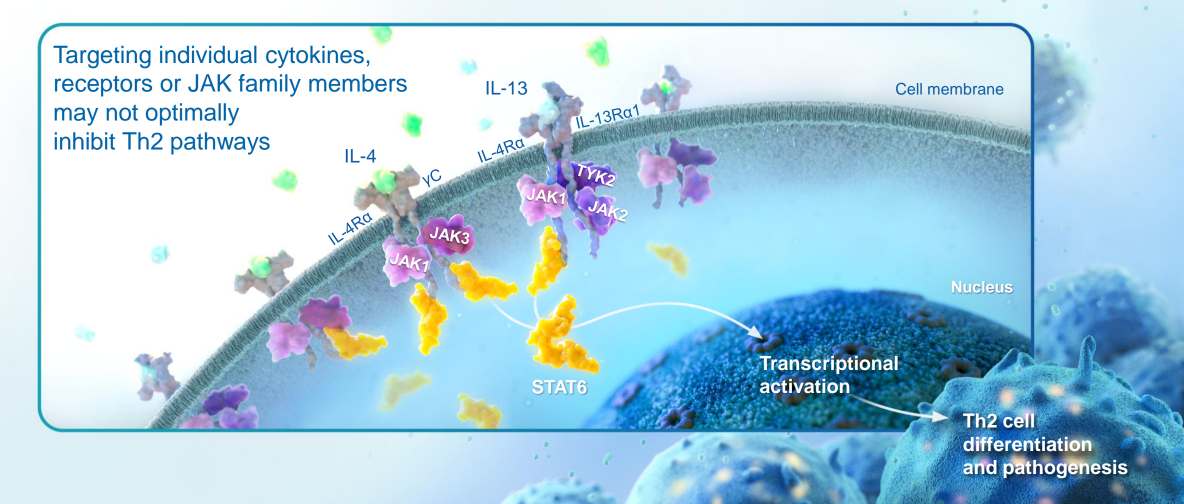
DIFFERENTIATED PRODUCT

Patient-friendly oral formulation may enable treatment beyond those currently served by injectable agents

Source: GlobalData 2023, ClearView HealthCare Partners

STAT6 is a First- and Best-In-Class Opportunity to Selectively Target Th2 Inflammatory Disease Pathways





STAT6 Inhibitor REX-8756 is Highly Potent and Selective in Biochemical and Cellular Assays



	Biochemical potency (SH2scan K _D)	Cellular potency (pSTAT6 IC ₅₀ in human PBMCs)	Biochemical STAT family selectivity	Cellular selectivity (PBMCs)	SH2 domain selectivity
REX-8756	0.04 nM	0.72 nM (IL-4) 0.19 nM (IL-13)	>1,000X vs. STAT1/2/3/4/5	>1,000X vs. STAT1/2/3/4/5	

STAT6 Inhibitor Demonstrates Biologic-Like Selectivity and Differentiates from JAK Inhibitors



Direct selective STAT6 inhibition provides greater selectivity than currently approved JAK inhibitors

T CELL FUNCTION

HEMATOLOGIC HOMEOSTASIS

		General adaptive immune response	Defense against viruses and bacteria	Defense against extracellular pathogens	Defense against parasites and mediation of antibody responses	Erythropoiesis	Thrombopoiesis
		T Cell Activation (CD25)	Th1 Cell Function (IFNγ)	Th17 Cell Function (IL-17A)	Th2 Cell Function (IL-5)	EPO-Induced STAT5-Driven Transcription	TPO-Induced STAT5-Driven Transcription
STAT6 Inhibitor	REX-8756	>3,000 nM	>1,000 nM	>2,500 nM	19 nM	>10,000 nM	>10,000 nM
IL-4 / IL-13 Antagonist	Dupilumab	>10,000 nM	>1,000 nM	>1,000 nM	26 nM	>1,000 nM	>1,000 nM
	Abrocitinib	1,300 nM	900 nM	73 nM	80nM	3,200 nM	2,800 nM
JAK Inhibitors	Upadacitinib	39 nM	36 nM	7.4 nM	4.3 nM	69 nM	20 nM
	Baricitinib	110 nM	210 nM	25 nM	15 nM	56 nM	42 nM

Recludix Pharma JPM Morgan Healthcare Conference 2023 Recludix Pharma data on file, March 2023

Selectivity relative to Th2 inhibition:

>30X

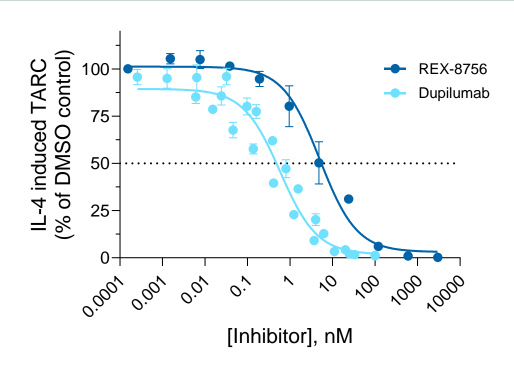
10-30X

<10X

STAT6 Inhibitor Fully Inhibits IL-4/13 Stimulated STAT6-Driven Biomarkers in Human PMBCs

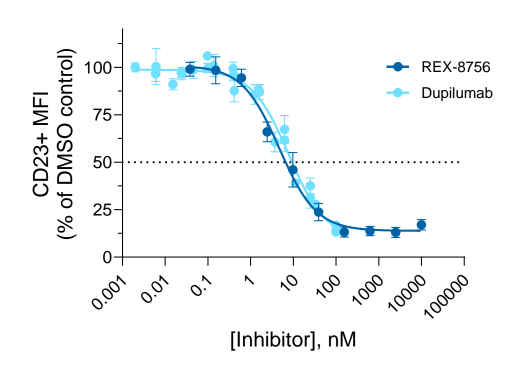


IL-4 induced TARC



	REX-8756	Dupilumab
IC ₅₀	6.3 nM	0.8 nM

IL-4 induced CD23 activation



	REX-8756	Dupilumab
IC ₅₀	5.1 nM	11 nM

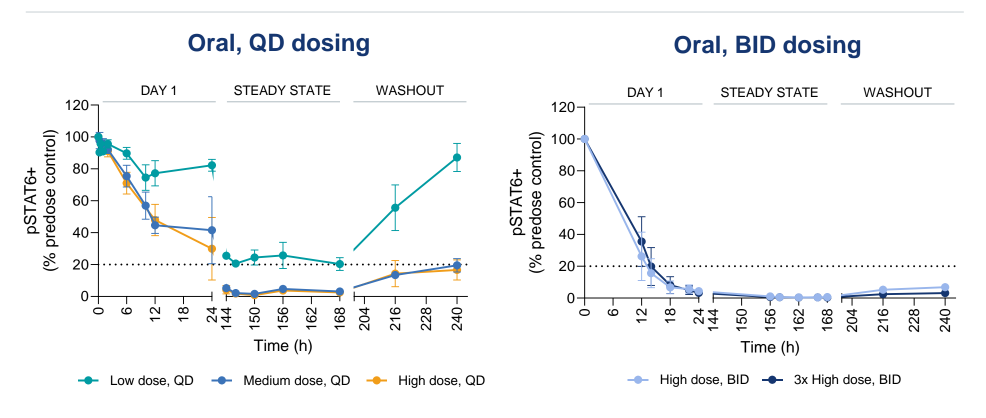
Recludix Pharma data on file, April 2022

Recludix Pharma

REX-8756 Achieves Complete pSTAT6 Inhibition with Once Daily Dosing and is Well Tolerated in Dog 7-Day Study



Efficacious dose at >80% pSTAT6 PD modulation



Multi-day PK/PD & Hazard ID study

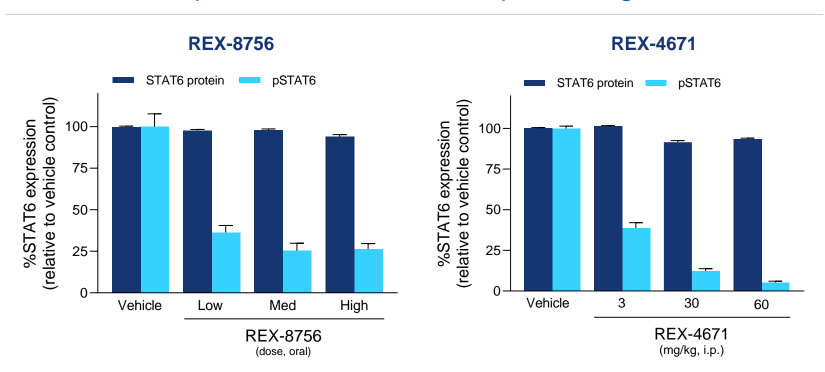
- Once or twice daily oral administration
 - Day 1
 - Day 7: Steady State
 - Day 9-10: Washout
- Clinical observations, hematology, serum chemistry, coagulation

- Highly selective pSTAT6 inhibition, no off-target pSTAT1 or pSTAT3 impact observed
- Well tolerated at all dose levels

Recludix STAT6 Inhibitors Achieve Deep and Durable pSTAT6 Inhibition In Vivo without STAT6 Protein Degradation in Mice

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- Recludix compounds achieve dose-dependent, rapid, and durable pSTAT6 inhibition in blood and tissues
- Reversible pSTAT6 inhibition achieves exquisite targeting of type-2 inflammation without STAT6 protein degradation

In vivo pSTAT6 inhibition without protein degradation



Single dose PK/PD study

- Samples collected 4hr post single dose
- Whole blood and splenocytes ex vivo stimulated with IL-4 to induce STAT6 pathway activation
- Total STAT6 and pSTAT6 quantified by flow cytometry

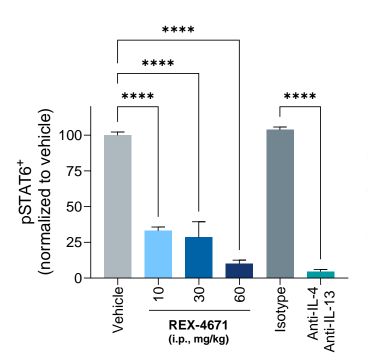
STAT6 Tool Compound REX-4671 Has Comparable Efficacy to Combined Anti-IL-4/13 Biologics in Dermatitis Model

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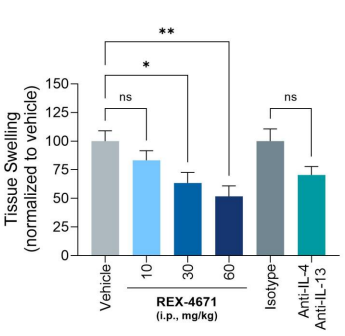
- REX-4671 dose dependently inhibits pSTAT6 and significantly reduced skin inflammation in chemicalinduced dermatitis model
- Comparable in vivo efficacy to the combination of anti-IL-4/13 surrogate antibodies

abdomenal ear ear inflammation FITC FITC SKIN SWELLING prophylactic prophylactic

Spleen pSTAT6 inhibition



Skin inflammation

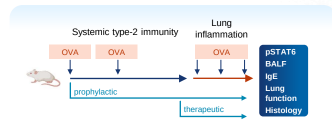


*p<0.05, **p<0.01, ***p<0.001, ****p<0.0001

Prophylactic REX-4671 Demonstrates Comparable Efficacy to Combined Anti-IL-4/13 Biologics in OVA-Asthma Model



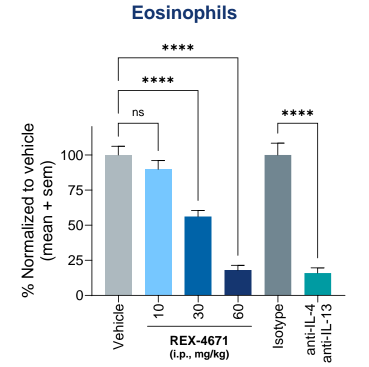
 REX-4671 demonstrates comparable efficacy to combined anti-IL-4/13 rodent surrogate antibodies



pSTAT6 inhibition

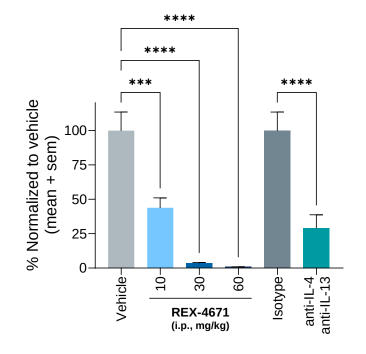
Whole Blood % Normalized to vehicle 125-100-(mean + sem) 75-25anti-IL-4 anti-IL-13 Vehicle-Isotype-**REX-4671** (i.p., mg/kg)

Lung immune cell infiltration



Humoral immunity

Serum OVA-specific IgE



Summary and Next Steps



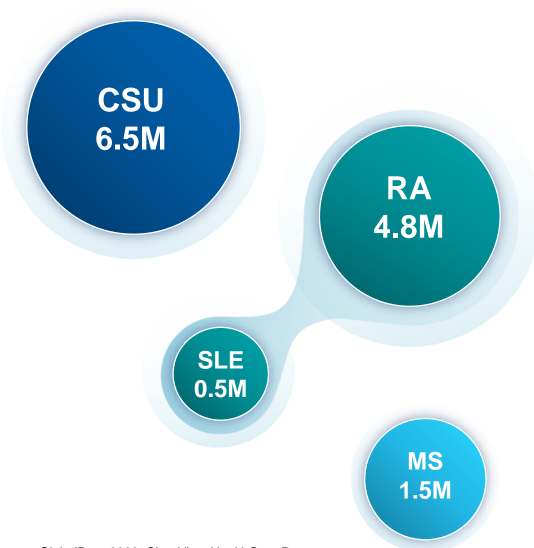
- REX-8756 is a potent and selective oral STAT6 inhibitor that demonstrates complete pathway inhibition and is well tolerated in preclinical studies
- Completion of GLP toxicology studies and Development Candidate nomination associated with a \$50 million payment to Recludix under the collaboration with Sanofi
- Investigational New Drug (IND) enabling activities ongoing to support IND submission 2H 2025



BTK

Selective Targeting BTK Has the Potential to Yield Superior Efficacy and Safety in Chronic Inflammatory Diseases





>13M
TOTAL PATIENT IMPACT

DIFFERENTIATED RELATIVE TO TRADITIONAL TKI APPROACH

Targeting SH2 domain leads to best-in-class selectivity to enable improved safety margins

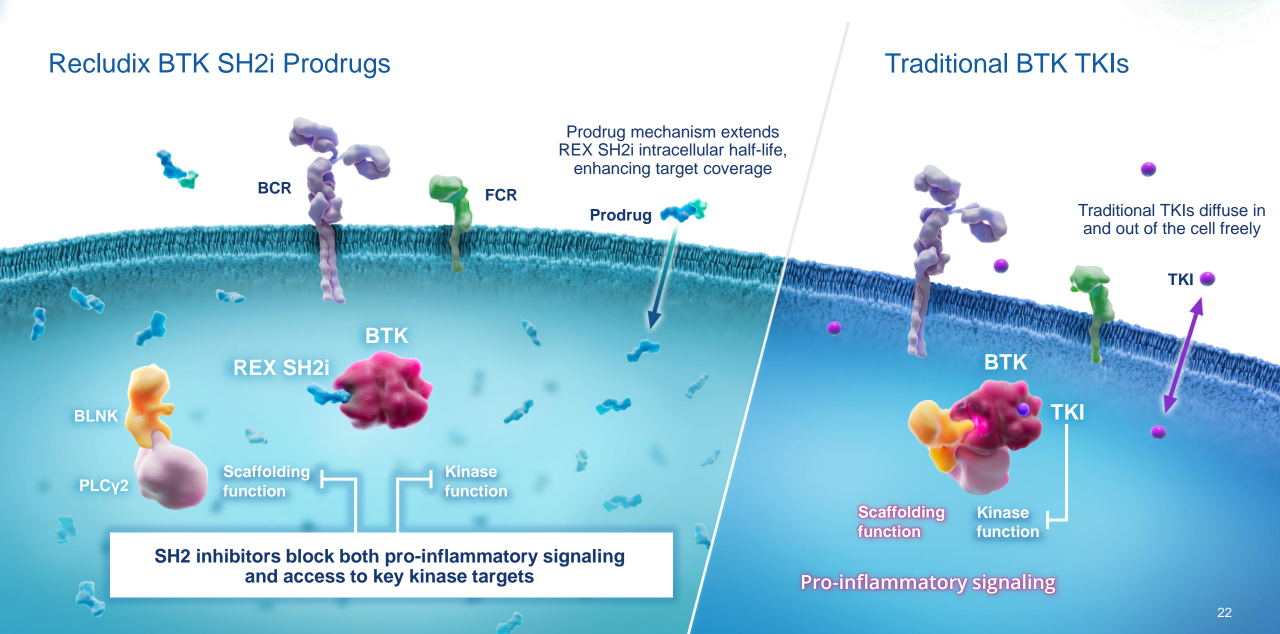
SH2 approach disrupts the central scaffolding function of BTK to deeply inhibit pro inflammatory signaling and widen the therapeutic window

Prodrug mechanism enhances target coverage to drive improved efficacy

Source: GlobalData 2023, ClearView HealthCare Partners

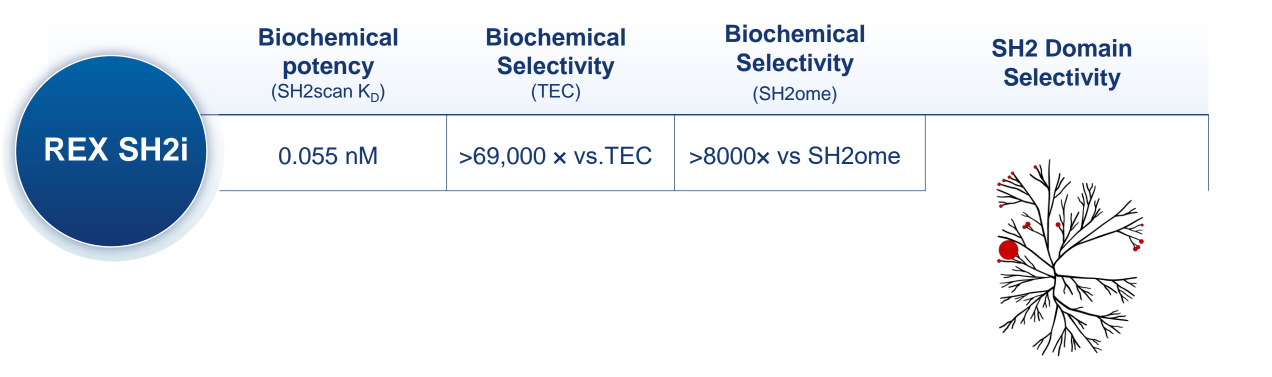
BTK SH2 Inhibitor is First-In-Class with Differentiated Profile Relative to Traditional TKIs





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Recludix Has Developed Highly Potent and Selective BTK SH2 Domain Inhibitors in Biochemical Assays



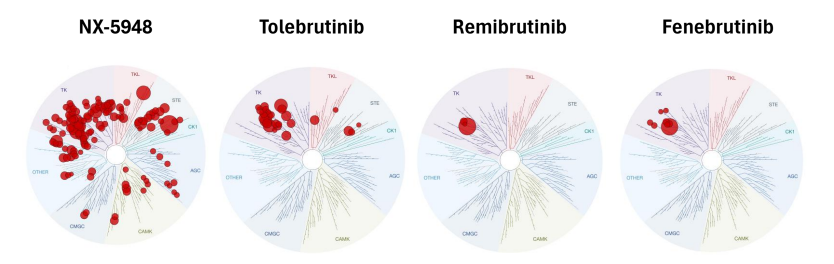
BTK SH2 Domain Inhibitors are Differentiated Relative to Traditional BTK Tyrosine Kinase Inhibitors



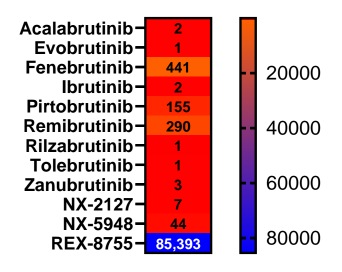
Selectivity difference is predicted to drive improved safety and efficacy outcomes

- Recludix BTK SH2 inhibitors:
 - Demonstrate exquisite selectivity
 - Do not inhibit TEC kinase, an important off-target involved in platelet dysfunction
 - Demonstrate TEC/BTK selectivity greater than 80,000x

Kinase selectivity



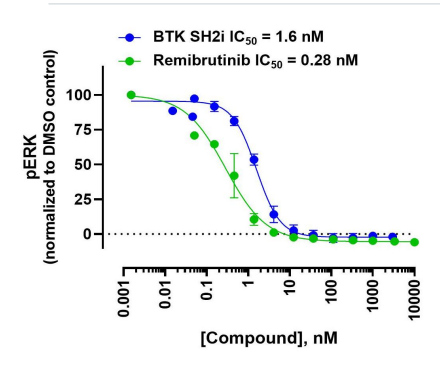
TEC/BTK selectivity



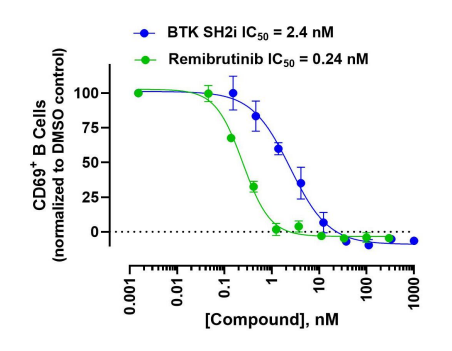
BTK SH2 Domain Inhibitors Fully Block pERK Signaling and Downstream Immune Cell Activation

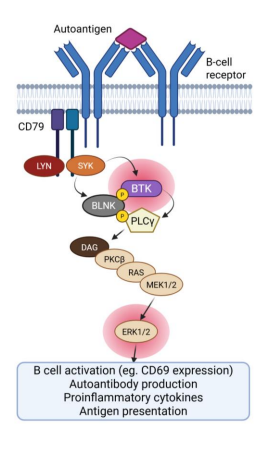


pERK Assay



CD69 Expression



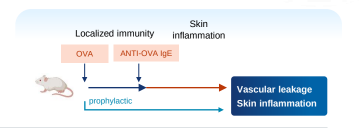


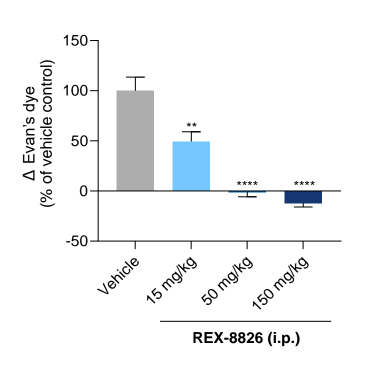
BTK SH2 Domain Inhibitors Abrogate Mast Cell Activation in In Vivo Model of Chronic Urticaria

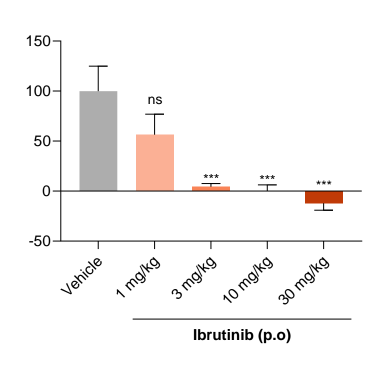


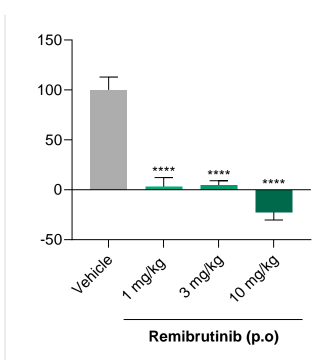
BTK SH2 domain inhibitors demonstrate strong dose-dependent efficacy

Vascular leakiness (Evan's dye extravasation)









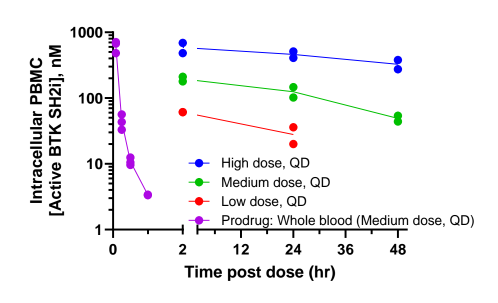
** p < 0.1, *** p < 0.001, **** p < 0.0001

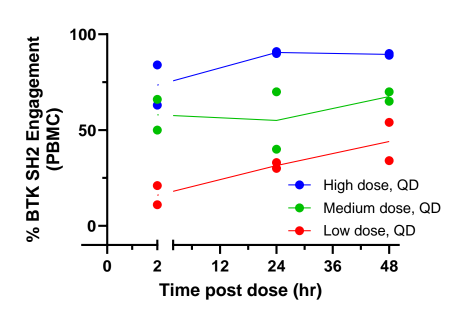
BTK SH2 Domain Inhibitors Achieve Durable Intracellular PK Leading to Prolonged Target Engagement in Dog PBMCs



BTK SH2i Pharmacokinetics

BTK Target Engagement





- Following intravenous dosing in dogs, active BTK SH2i maintained durable intracellular exposures in target PBMCs while exposure of the prodrug was transient
- BTK SH2i exposure in PBMCs enabled deep, durable, and dose-dependent target engagement



Highlights and Upcoming Milestones

Highlights and Near-Term Milestones





KEY ACCOMPLISHMENTS

- Successfully drugged previously "undruggable" SH2 domains
- Advanced STAT6 inhibitor in global partnership with Sanofi
 - Potent, selective, reversable, and orally bioavailable compounds
 - Favorable differentiation from IL-4/IL-13 biologics and JAK/TYK2 inhibitors
 - In vivo efficacy and target modulation, without protein degradation, in inflammation disease models
 - GLP toxicology studies completed and Drug Candidate nominated, triggering \$50 million milestone payment from Sanofi



NEAR-TERM MILESTONES

STAT6

- IND submission 2H25
- Phase 1 study initiation 2H25

BTK

 Continue to advance program towards clinic



Thank you

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Unlocking New Therapeutic Possibilities