September 2022 Innovating for Everyone

Clinical-stage biopharmaceutical company focused on next generation therapeutics meeting unmet patient needs.



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Our Mission



At Hoth Therapeutics, we strive to develop innovative, impactful, and ground-breaking treatments with a goal to improve patient quality of life. We are a catalyst in early-stage pharmaceutical research and development, elevating promising drugs from the bench to pre-clinical and clinical testing. Utilizing a patient-centric approach, we collaborate and partner with a team of scientists, clinicians, and key opinion leaders to seek out and investigate medications that hold immense potential to create breakthroughs and diversify treatment options. Our mission is to bring value to both our shareholders and our patient populations.



Key Investment Highlights





Clinical Programs



Robust Pre-Clinical Development Programs

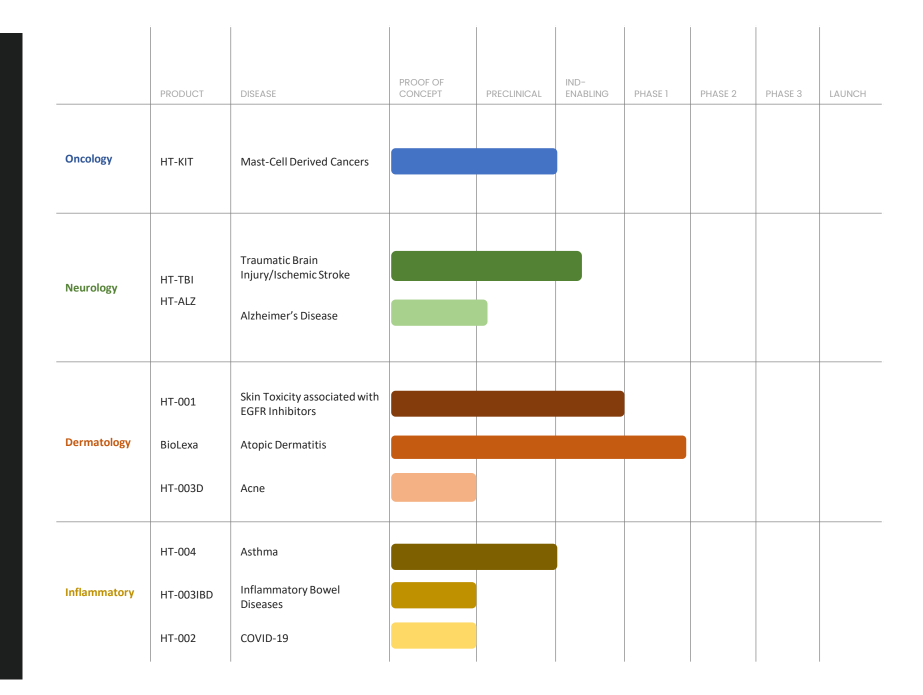


Targeting Unmet Medical Needs to Address Broad Market



Experienced Management and Advisory Board

Pipeline: Multiple Shots on Goal





Primary Development



HT-001 Topical Gel

HT-KIT Injection

HT-TBI Suspension IM

HT-ALZ Oral Soluble Film

BioLexa Lotion

HT-001: Value Proposition



Market Growth: EGFR Inhibitor Skin Toxicity market predicted to grow from \$52M in 2018 to \$391M by end of 2030*

Mechanism of Action: 12-week study conducted at GW suggests the topical application of HT-001 significantly reduces erlotinib-induced cutaneous toxicities applied preventatively or proactively. It supports that HT-001 may be used as a topical intervention to treat EGFR-inhibitor-induced cutaneous toxicity.**

Addresses Unmet Need: No current approved product on the market that specifically treats EGFR inhibitor cutaneous toxicities, which occur in up to 90% of patients undergoing EFGR inhibitor therapy.***

*EGFR Inhibitors-Induced Skin Disorders-Market Insights, Epidemiology, and Market Forecast-2030

**https://ir.hoththerapeutics.com/ht-001

***https://jamanetwork.com/journals/jamadermatology/article-abstract/2767656

Recent & Upcoming Milestones: HT-001 Topical Gel



LogicalLogical2022Jan 2022IND-Enabling studies
completed in Q1 2022Proactive treatment of
EGFRI skin toxicity
demonstrated in animal

model

Feb 2022

Selection of GMP Manufacturer – Nuvisan Pharma February 2022



2023

IND Submission/Clinical Trial target for Q1 2023

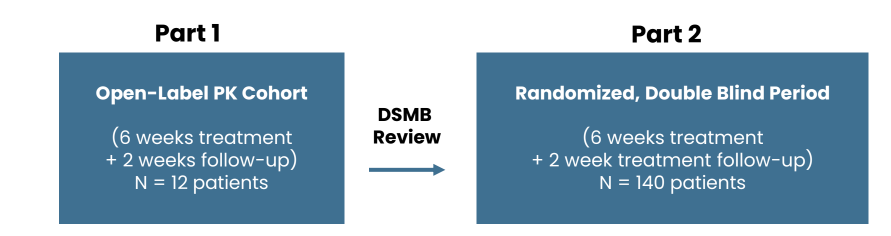
IND-Opening Trial: CLEER-001 Phase 2a Dose Ranging Study



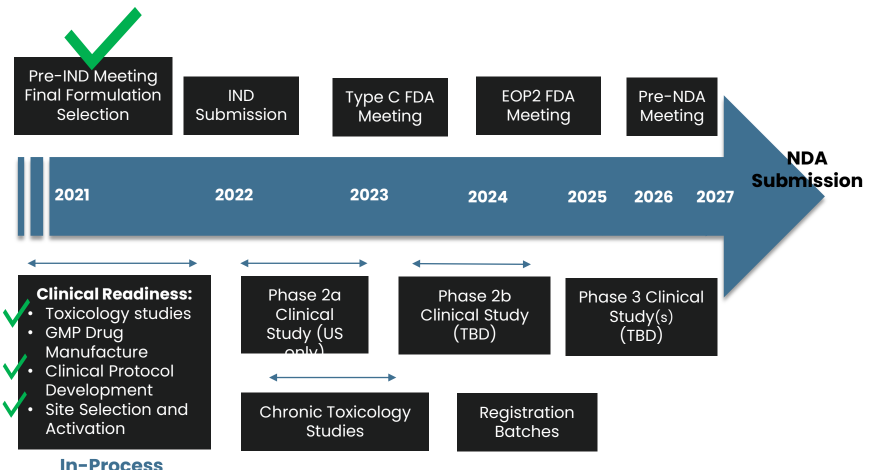
A Randomized, Placebo-Controlled, Parallel Phase 2a Dose Ranging Study to Investigate the Efficacy, Safety, and Tolerability of Topical HT-001 for the Treatment of Cutaneous Toxicities Associated with EGFR Inhibitors

2 Parts:

- Part 1: Open-label treatment with HT-001 2% for 6 weeks (PK cohort)
- Part 2: Randomized double-blind treatment with HT-001 0.5%, 1%, 2% or placebo



HT-001 **505**(b)(2) **Development** Pathway



In-Process



Current estimated dates; pending FDA meetings for phase 2b/phase 3 clinical studies

Erlotinib (noVeh C) – Week 12

Proactive **Topical HT-**001 Significantly **Reduces EGFR** Inhibitor-Induced Cutaneous Toxicity





Erlotinib + Topical HT-001 Initiated Week 1 – Week 12



Erlotinib + Topical HT-001 Initiated Week 6 - Week 12



| Group | Change Compared to Erlotinib Only Group: Facial Skin Lesions at 12 weeks | Change Compared to Erlotinib Only Group: Hair Loss at 12 weeks |
|---|--|---|
| Preventati ve Topical HT-001 + Erlotinib | 58.5% Reduction (p<001 vs Erl and p<0.01 vs control) | 56.2% reduction in hair loss (p<0.001 vs Erl and p<0.001 vs control) |
| Proactive (week 6) Topical HT-001 + Erlotinib | 47.8% reduction (p<0.001 vs Erl and p<0.001 vs control) | 44.4% reduction in hair loss (p<0.001 vs Erl and p<0.001 vs control) |

HT-KIT: Value Proposition



Market Growth: Global systemic mastocytosis treatment revenue is \$128M and projected to grow at 5.8% CAGR through 2031*

Mechanism of Action: HT-KIT is an antisense oligonucleotide that results in non-functional cKIT via mRNA frameshift.**

Addresses Unmet Need: KIT D816V mutation found in >80% of adult systemic mastocytosis cases results in confirmational changes that make some tyrosine kinase inhibitor drugs ineffective.**

cKIT is also implicated in gastrointestinal stromal tumors, acute myeloid leukemia, and other rare cancers

*Global Systemic Mastocytosis Treatment Market Research Report, January 2022, Market.US

**Snider et al., Targeting KIT by frameshifting mRNA transcripts as a therapeutic strategy for aggressive mast cell neoplasms, Molecular Therapy (2021), https://doi.org/10.1016/j.ymthe.2021.08.009

Recent & Upcoming Milestones: HT-KIT Injection





Jan 2022

Preclinical proof of concept studies initiated with NC State



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Mar 2022

FDA Grants Orphan Drug Designation



Q4 2022

API Manufacturing

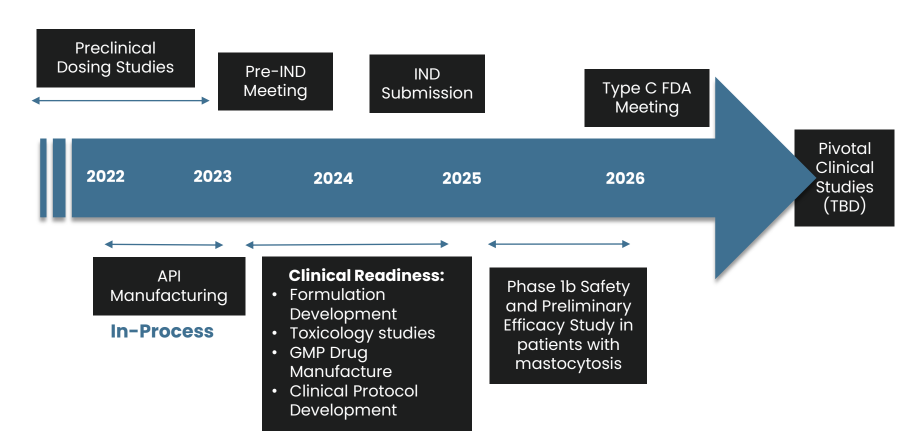
IND-enabling PK/PD animal model studies

List

2023

Pre-IND Submission target for 2023 Formulation Development HT-KIT Orphan Drug Development Pathway





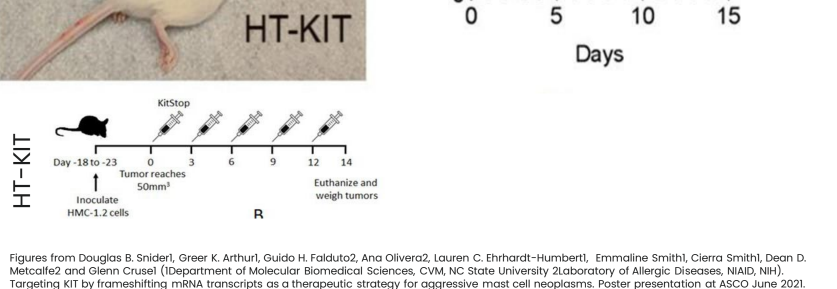
Current estimated dates; pending FDA meetings for clinical studies.

Systemic delivery of human HT-KIT inhibits tumor growth in a humanized xenograft mast cell neoplasia model

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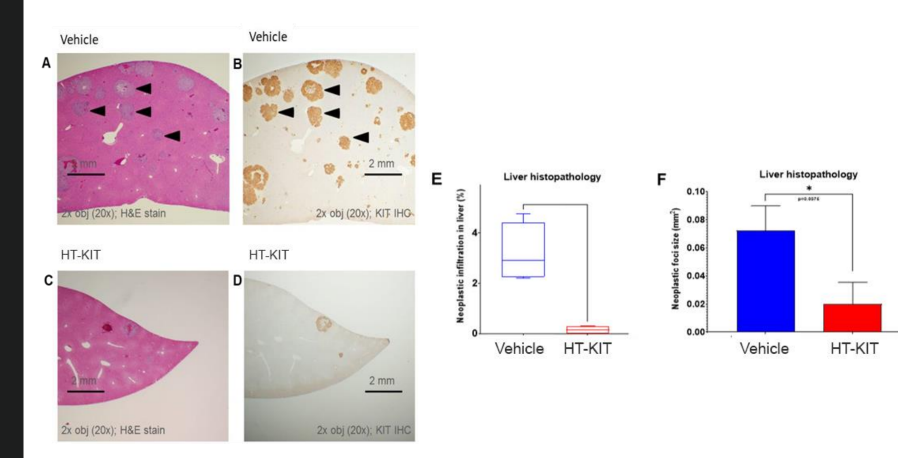
Vehicle





HMC-1.2 cells В Vehicle Tumor volume (mm³) 200 200 HT-KIT ****

HT-KIT Reduces Liver Infiltration of Neoplastic Mast Cells in a Humanized Xenograft Model of Mast Cell Neoplasia





Figures from Douglas B. Sniderl, Greer K. Arthurl, Guido H. Falduto2, Ana Olivera2, Lauren C. Ehrhardt-Humbertl, Emmaline Smithl, Cierra Smithl, Dean D. Metcalfe2 and Glenn Crusel (IDepartment of Molecular Biomedical Sciences, CVM, NC State University 2Laboratory of Allergic Diseases, NIAID, NIH). Targeting KIT by frameshifting mRNA transcripts as a therapeutic strategy for aggressive mast cell neoplasms. Poster presentation at ASCO June 2021.

HT-TBI: Value Proposition



Market Growth: The global stroke management market size was valued at \$31.7B in 2020, and is projected to reach \$67.7B by 2030, registering a CAGR of 7.8% from 2021 to 2030.*

The traumatic brain injuries treatment market to account \$201.1B by 2029 by growing at a CAGR of 5.00% in the forecast period of 2022-2029.**

Mechanism of Action: Target neurokinin-1/substance P to prevent downstream inflammatory pathways leading to secondary brain injury (edema, inflammation)

Addresses Unmet Need: : There is insufficient clinical evidence to support the use of the current hyperosmotic therapy methods (eg, mannitol) for lowering intracranial pressure per the Guidelines for the Management of Severe Traumatic Brain Injury; hyperosmotic therapy can also be associated with risk to patient morbidity and mortality.***

*Stroke Management Market Statistics 2030, Allied Market Research

**Global Traumatic Brain Injuries Treatment Market – Industry Trends and Forecast to 2029, Data Bridge Market Research

***Guidelines for the Management of Severe Traumatic Brain Injury 4th Edition, Brain Trauma Foundation

Upcoming Milestones: HT-TBI Suspension IM



Q3 2022

Begin formulation development and feasibility batches for upcoming Pre-IND submission. Q4 2022 Injection device evaluation and feasibility

testing

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Q1 2023

GMP manufacturing target

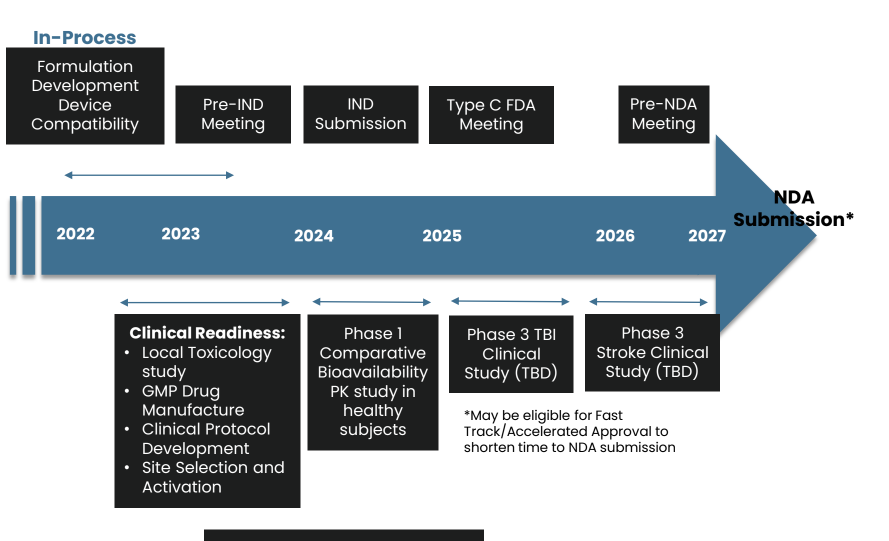
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Q2 2023

Pre-IND Meeting with FDA target

HT-TBI 505(b)(2) Development Pathway





Registration Batches

Current estimated dates; pending FDA meetings for phase 3 clinical studies.

HT-ALZ: Value Proposition



Market Growth: The Global Alzheimer's Disease Treatment Market is expected to grow at a CAGR of around 12.8% from 2020 to 2027 and reach the market value of over \$5.2B by 2027.*

Mechanism of Action: HT-ALZ targets Substance P** in the brain, which has both negative (inflammatory) and positive (anti-amyloidogenic, memory, neuroprotective) roles in Alzheimer's disease. The precise MoA of HT-ALZ is currently under investigation.

Addresses Unmet Need: : There are currently no drugs approved that are considered disease modifying and demonstrate cognitive improvement. Preclinical data with HT-ALZ indicates HT-ALZ may provide both therapeutic benefits.

*https://www.acumenresearchandconsulting.com/alzheimers-disease-treatment-market

**Martinez AN, Philipp MT. Substance P and Antagonists of the Neurokinin-1 Receptor in Neuroinflammation Associated with Infectious and Neurodegenerative Diseases of the Central Nervous System. J Neurol Neuromedicine. 2016;1(2):29-36. doi:10.29245/2572.942x/2016/2.1020

**Severini C, Petrella C, Calissano P. Substance P and Alzheimer's Disease: Emerging Novel Roles. Curr Alzheimer Res. 2016;13(9):964-72. doi: 10.2174/1567205013666160401114039. PMID: 27033058.

Recent & Upcoming Milestones: HT-ALZ Oral Soluble Film





2021

Initiated Proof-of-Concept study with Alzheimer's Disease leader, Washington University ("WashU")

Jan 2022

Initial data from WashU studies show a significant decrease in brain interstitial fluid Aβ in an Alzheimer's disease mouse model

July 2022

Chronic treatment with HT-ALZ (>5 weeks) demonstrated significant improvement in behavioral and cognitive function tests in an Alzheimer's disease (AD) mouse model

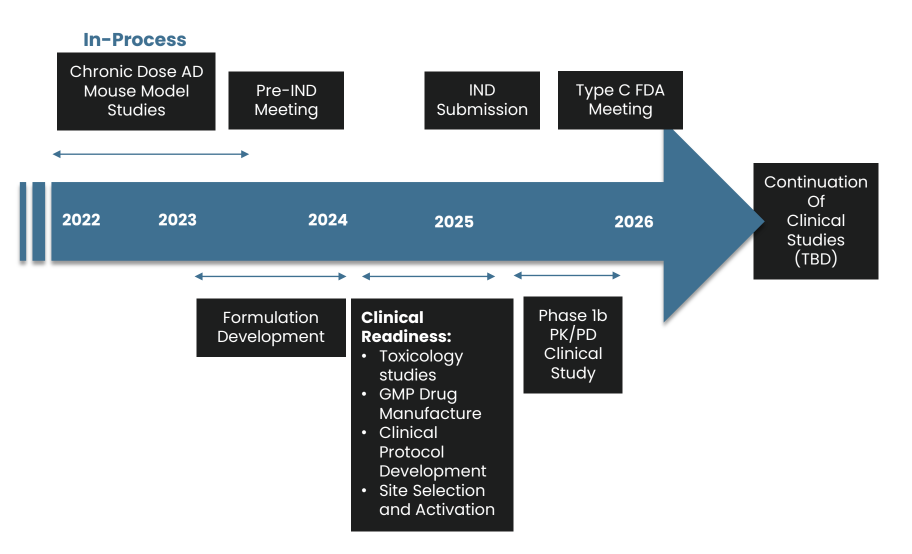
Additional chronic dose ranging studies initiated



2023

Oral formulation development target initiation date

HT-ALZ 505(b)(2) Development Pathway





Current estimated dates; pending FDA meetings for clinical studies.

Biolexa: Value Proposition

Market Growth: Atopic dermatitis market predicted to grow from \$6.4B in 2017 to \$18.3B by end of 2027*

Mechanism of Action: Novel mixture of two previously approved compounds targeting the underlying Staphylococcus aureus infection hypothesize to potentiate Atopic Dermatitis (AD) or eczema flares – First compound prevents biofilm formation, which protects the underlying infection, allowing the second, an antibiotic, to more effectively treat the underlying infection.

Addresses Unmet Need: Non-corticosteroid treatment targeted for treatment of both pediatric and adult mild to moderate AD populations

*Atopic Dermatitis Market – Global Industry Analysis, Size and Forecast, 2017-2027



Recent & Upcoming Milestones: BioLexa Lotion





2021 Phase 1b Cohort 1 with

healthy subjects

completed

Dec 2021 Phase 1b cohort

Phase 1b cohort in patients with mild to moderate atopic dermatitis initiated



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Sep 2022

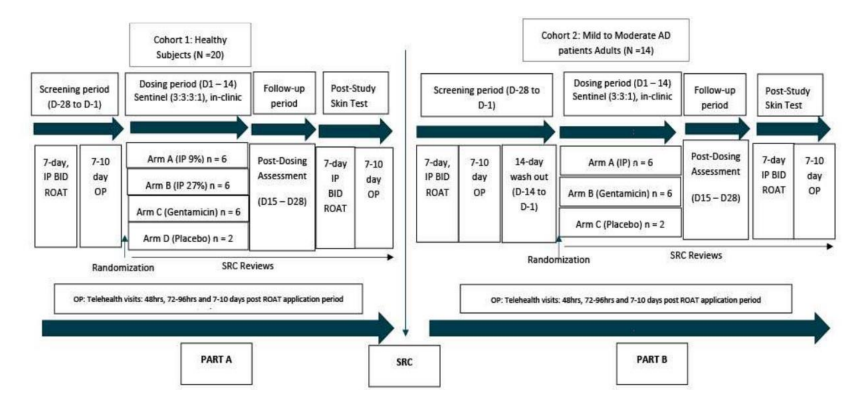
Open Label dosing portion of Phase 1b Study is completed



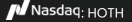
2023

Data Readout from Phase 1b Clinical Study

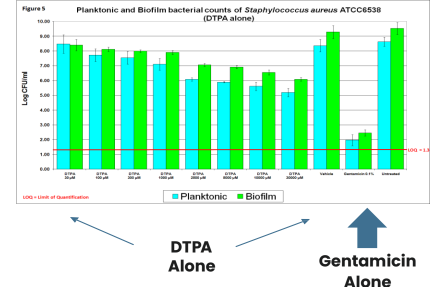
BioLexa Phase 1b Clinical Study Design A Randomised, Double-Blind, Vehicle Controlled, Sequential Group Study to Determine the Safety, Tolerability, Pharmacokinetics and Efficacy of Twice Daily Application of Topical BioLexa[™] in Adult Healthy Subjects and Patients with Mild to Moderate Atopic Dermatitis

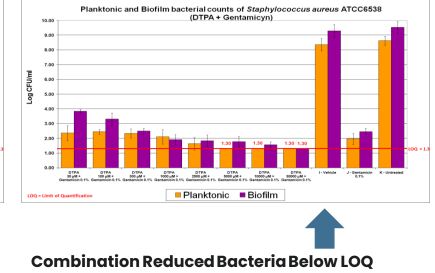






BioLexa: Proof-of-Concept Results This study concluded that the combination of gentamicin and Ca-DTPA is more effective to reduce bacteria growth and inhibit the formation of biofilms than each compound individually.





Miller School of Medicine, of the University of Miami and University of Cincinnati – Determination of the effects of a novel antimicrobial agent used in conjunction with Gentamicin on *Staphylococcus aureus* using a porcine model: preliminary evaluations Jose Valdes, Joel Gil, Andrew Herr, Andrew Harding and Stephen Davis Miller School of Medicine, of the University of Miami and University of Cincinnati – Determination of the effects of a novel antimicrobial agent used in conjunction with Gentamicin on *Staphylococcus aureus* using a porcine model: preliminary evaluations Jose Valdes, Joel Gil, Andrew Herr, Andrew Harding and Stephen Davis



Preclinical Development



HT-004

Direct Detect Breath Diagnostic Device

HT-003: Value Proposition



Market Growth: The global acne treatment market size was valued at \$5.46 Bn in 2019 & is projected to reach \$7.19 Bn by 2027, with a CAGR of 4.6% during forecast period.* IBD Treatment market was valued at USD 15.87 billion by 2019, growing with 4.1% CAGR during the forecast period, 2020-2026**

Mechanism of Action: Group of novel inhibitors of retinoic acid metabolism (collectively called RAMBAs), which prolong the presence of retinoic acid. Retinoids play key role in the regulation of immune cells and inflammation and are also important for differentiation of the skin.

Addresses Unmet Need: Focuses on restoring immune system rather than inducing immune suppression (current therapies)

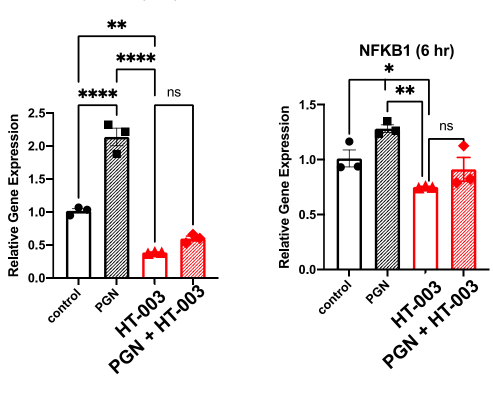
*https://www.fortunebusinessinsights.com/acne-treatment-market-103361

**https://www.marketwatch.com/press-release/inflammatory-bowel-disease-ibd-treatment-market-analysis-share-trends-size-forecast-from-2020---2030-2021-11-01?tesla=y

HT-003D: Dermal Preclinical Study Results



TLR2 (6 hr)





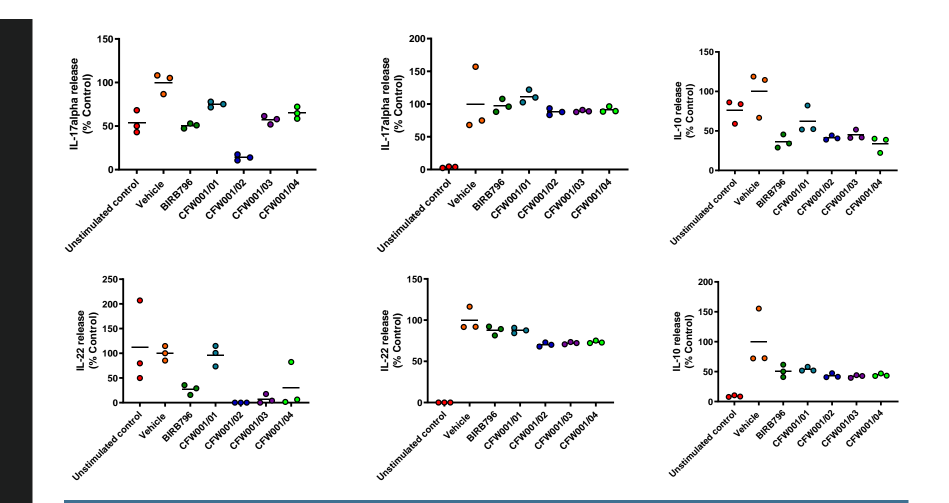
TLR2 is one of the most critical genes for acne pathophysiology



Data shows that HT-003 significantly downregulates TLR2 expression after challenge with PGN (TLR2 agonist) in an in vitro human keratinocyte model

HT-003IBD: Preclinical Study Results Ulcerative Colitis Ex Vivo Tissue (n = 2 donors)



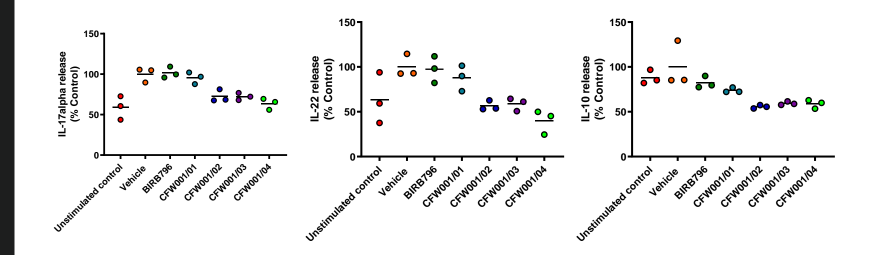


HT-003 molecules reduce inflammatory cytokines associated with IBD and promotes intestinal homeostasis

BIRB796 = positive therapeutic control | CFW = HT-003 molecules 1 – 4 unique entities

HT-003IBD: Preclinical Study Results Crohn's Ex Vivo Tissue (n = 1 donor)





HT-003 molecules reduce inflammatory cytokines associated with IBD and promotes intestinal homeostasis

CFW = HT-003 molecules 1 – 4 unique entities | BIRB796 = positive therapeutic control

HT-004: Value Proposition

Market Growth: The global asthma therapeutics market reached a value of \$17.6B in 2020. The market is expected to reach a value of \$19.13B by 2026, expanding at a CAGR of 1.60% during 2021-2026.*

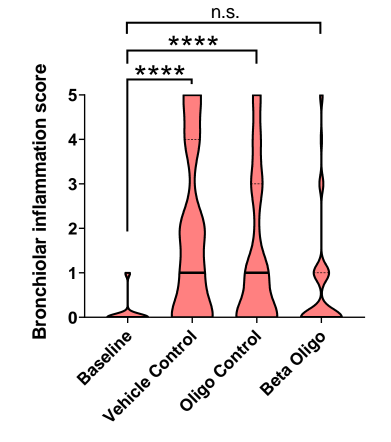
Mechanism of Action: Target IgE receptor trafficking to prevent downstream inflammatory pathways

Addresses Unmet Need: New class of drug for maintenance treatment of asthma with potential for a better safety profile

*Asthma Therapeutics Market: Global Industry Trends, Share, Size, Growth, Opportunity and Forecast 2021-2026, imarc



HT-004: Asthma & Allergic Inflammation



- Peribronchiolar Inflammation was reduced by inhalation of HT-004 that targets FcER1beta alternative exon splicing.
- Ovalbumin inhalation induced airwaycentric recruitment of inflammatory cells predominated by eosinophils admixed with lymphocytes, macrophages, and fewer mast cells.
- Inflammatory cell recruitment was minimal in lungs of mice lacking the ovalbumininduced allergic airway disease and administered only PBS vehicle control.
- Inflammatory cell recruitment was moderate to marked resulting in expansion of peribronchiolar connective tissues by several cells thick in some areas for mice in control treatment groups with ovalbumininduced allergic airway disease (vehicle control and oligo (non-target) control.
- Despite ovalbumin-induced allergic airway induction, lungs from mice receiving inhalation of HT-004 had reduced inflammatory cell recruitment around bronchioles.



Breath Detect Diagnostic Device



Sensor on Chip нотн Breath Collection Tube

Direct Detect Breath Diagnostic Device System: Value Proposition



- Technology can be developed for home use by patients or lay users for rapid, large-scale screening
- ✓ Breath samples can be collected easily and noninvasively without additional sample collection devices or sample processing
- ✓ Results available in a time scale of minutes using smart phone camera or handheld, multi-use detectors (to be designed)
- Device has the potential to detect a broad range of target analytes with low limit of detection
- Sensor has been designed for large scale manufacturing with easily modifiable options for different targets/indications

Market Overview:

Attractive Opportunities in the Point of Care Market





Market growth in Latin America can be attributed to government initiatives, rising awareness about self-testing products, rapid expansion of the healthcare infrastructure, and increase in collaborations among key players in the point-of-care diagnostics market in the region.



CAGR of 10.8%

43.2

The global point-of-care diagnostics market is projected to reach USD 72.0 billion by 2027, growing at a CAGR of 10.8% during the forecast period.



The rising prevalence of infectious diseases and target conditions, the shortage of technicians to conduct conventional lab-based tests, supportive government policies, and the rising number of CLIA approvals for POC tests are the key factors driving the point-ofcare diagnostics market.

USD Billion

2022-е



The North American point-of-care diagnostics market is projected to reach USD 29.12 billion by 2027, growing at a CAGR of 11.3% during the forecast period.



Emerging markets, healthcare decentralization, and growing investments and funding for product development are expected to offer growth opportunities for market players during the forecast period.

USD Billion

2027-p



Latin America is the fastestgrowing regional market for point-of-care diagnostic products, with Brazil and Mexico emerging as major growth hotspots.

Point of Care & Rapid Diagnostics Market by Product (Glucose, Infectious Disease(HIV, Hepatitis C), Pregnancy Test), Platform (Microfluidics, Dipsticks), Mode of Purchase (OTC, Prescription), End User (Pharmacy, Hospital, Home) - Global Forecast to 2027. Markets and Markets.

Investment Highlights

Two Programs in Clinical Stage of Development

Diverse and Robust Pipeline of Pre-Clinical Candidates

- Addressing multi-billion-dollar unmet market opportunities across indications
- BioLexa Lotion Novel mixture of two FDA-approved compounds in clinical phase of development
- HT-001 no approved product/competitor currently on the market, clinical trial projected for early 2023

- Offers strong intellectual property portfolio, including exclusive licenses to patents and trademarks
- Multiple shots on goal with diversified portfolio and market
- Multiple assets have platform technology possibilities

Clean Financials

- 32.2 million shares outstanding (as of September 13, 2022)
- Cash on hand is sufficient to take company through the clinical and pre-clinical programs in current pipeline

Experienced Management, Board and Scientific Advisors

Experienced management team, board of directors and scientific advisors with proven financial, capital markets and drug development experience



Management Team



Robb Knie Chief Executive Officer



Stefanie Johns, Ph.D. Chief Scientific Officer



David Briones Chief Financial Officer



Hayley Springer Executive Vice President, Operations



Thank You.



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