



# **RedHill Biopharma Ltd.**

**("RDHL")**

Corporate Presentation  
August 2021

# Forward Looking Statement

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All forward-looking statements included in this presentation are made only as of the date of this presentation. We assume no obligation to update any written or oral forward-looking statement made by us or on our behalf as a result of new information, future events or other factors.

## Corporate Highlights

An emerging U.S. specialty biopharmaceutical company (Nasdaq: RDHL), primarily focused on U.S. commercialization and development of drugs for gastrointestinal (GI) diseases and infectious diseases

### Strong U.S. Commercial Footprint and Robust Development Pipeline with Multiple Near-Term Milestones

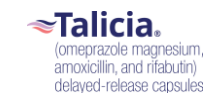
Promoting Three  
FDA-Approved Drugs

Multiple Phase 3 and  
Phase 2 Programs



# Emerging U.S. Specialty Pharma: Select Programs<sup>i</sup>

## Commercial Products<sup>ii</sup>



**Talicia® (omeprazole magnesium, amoxicillin and rifabutin) - *H. pylori* infection in adults**

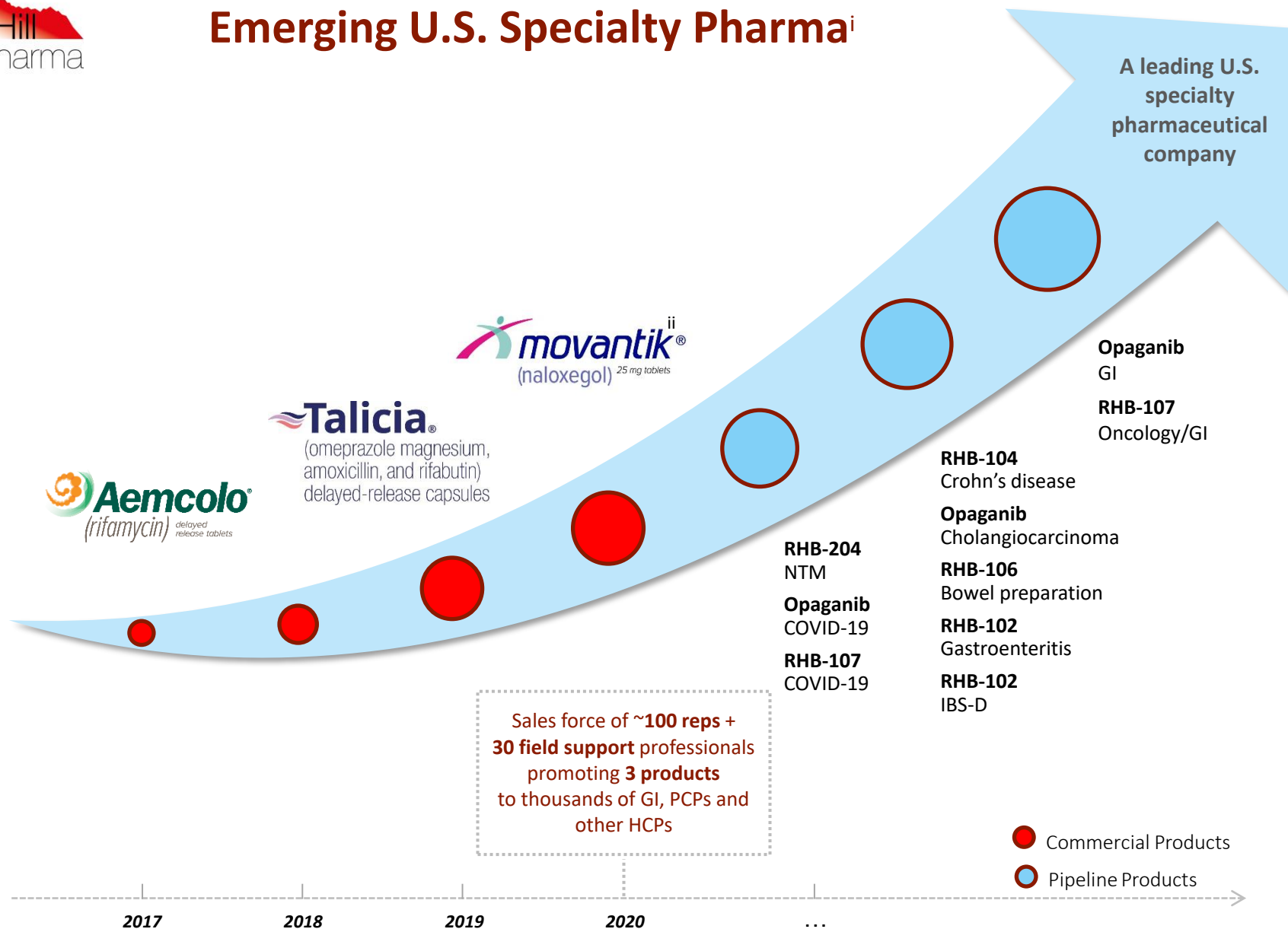
**Movantik® (naloxegol) - Opioid induced constipation (OIC) in adults with chronic non-cancer pain<sup>iii</sup>**

**Aemcolo® (rifamycin) - Travelers' diarrhea caused by noninvasive strains of *E. coli* in adults**

Development Pipeline <sup>iv</sup>		Pre-Clinical	Phase 1/2	Phase 3	NDA
RHB-204	NTM disease	Phase 3 U.S. study ongoing			
RHB-104	Crohn's disease	Positive results from Phase 3 MAP US study			
RHB-102	Gastroenteritis	Positive results from Phase 3 U.S. study			
	IBS-D	Positive results from Phase 2 U.S. study			
RHB-106	Bowel cleanser	Phase 2/3 studies planned			
Opaganib	Oncology Indications + COVID-19	Ongoing Phase 2/3 COVID-19 & Phase 2 oncology program			
RHB-107 (upamostat)	Oncology/GI + COVID-19	Ongoing Phase 2/3 COVID-19, GI & oncology indications			

<sup>i</sup> Estimated timeline/indication in the pipeline is subject to changes in development plans and regulatory requirements/clarifications, including complementary/additional studies; <sup>ii</sup> For full prescribing information see: Aemcolo®: [www.aemcolo.com](http://www.aemcolo.com); Talicia®: [www.talicia.com](http://www.talicia.com); Movantik®: [www.movantik.com](http://www.movantik.com); <sup>iii</sup> Movantik® is a registered trademark of AstraZeneca; <sup>iv</sup> Bekinda® (RHB-102) and Yeliva® (opaganib) are proposed tradenames which are subject to FDA review and approval at the time of NDA filing;

# Emerging U.S. Specialty Pharma<sup>i</sup>



<sup>i</sup> Presented events are forward looking statements and estimates and are subject to uncertainties including, among others, clinical and regulatory outcomes, marketing approvals, financial resources and commercial viability; This slide and strategic plan is made for illustrative purposes only. Please see "Disclaimer and Forward Looking Statements".

<sup>ii</sup> Movantik® is a registered trademark of AstraZeneca.

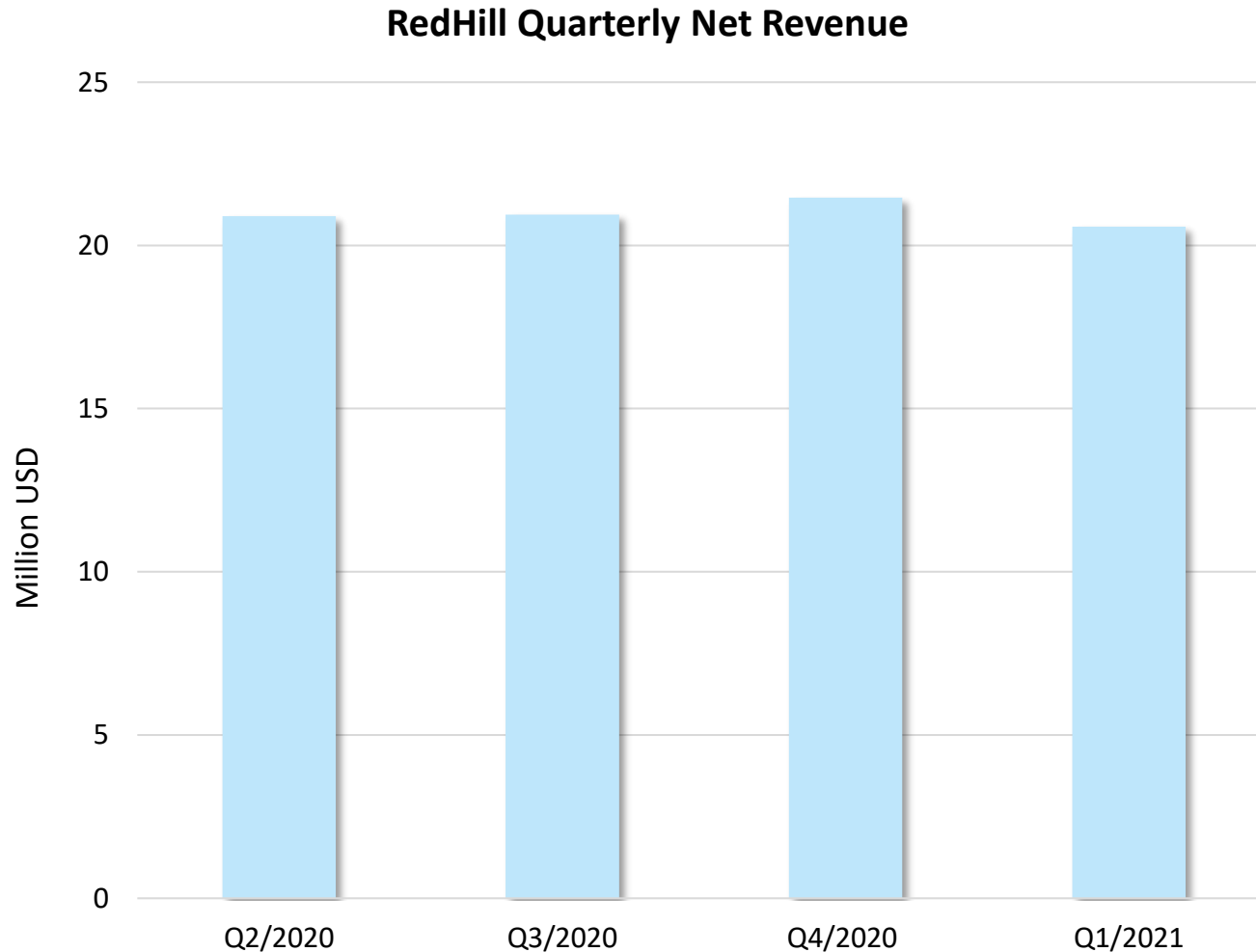
# Financial Highlights<sup>i</sup>

## RedHill Biopharma Ltd. Nasdaq: RDHL

Market Cap (approx.)	\$333 million
American Depositary Shares (ADSs)	46.67 million (Representing 466.7 million ordinary shares outstanding)
Cash Balance as of March 31, 2021 <sup>ii</sup>	Approx. \$92 million

<sup>i</sup> Financial information as of August 6, 2021 unless otherwise noted <sup>ii</sup> cash, cash equivalent, short term deposits and restricted cash.

# Revenue Growth Despite Challenging Pandemic Conditions



# Highly Experienced Senior Leadership

**Dror Ben-Asher**, Chief Executive Officer  
P.C.M.I. Ltd.

**Adi Frish**, Chief Corporate & Business Development Officer  
Y. Ben-Dror, MediGus

**Micha Ben Chorin**, Chief Financial Officer  
GVT, Pyramid Analytics, Starhome B.V.

**Reza Fathi, PhD**, Senior VP R&D  
XTL, PharmaGenics, Harvard Inst. of Chem. & Cell Biology

**David Wasserman**, Senior VP Alliance & Project Management  
Salix, Watson Pharmaceuticals, Glaxo PLC

**Rob Jackson**, Senior VP Sales & Marketing  
Salix, Vicuron, Merck & Co.

**Patricia Anderson**, Senior VP Regulatory Affairs  
MAPI Group, OptumInsight, Bayer, Novopharm

**Steven Thomasian**, VP Supply Chain  
Salix, Kala Pharmaceuticals, Cempira Pharmaceuticals

**Aida Bibliowicz**, VP Clinical Affairs Europe  
MSc Technion, MBA TAU, Cato Research Israel

**Todd Krzyzaniak**, VP Finance US Operations  
Salix Pharmaceuticals, BioDelivery Sciences International

**Gilead Raday**, Chief Operating Officer  
MSc Neurology, MBE Cambridge, Sepal Pharma

**Guy Goldberg**, Chief Business Officer  
Eagle Pharma, ProQuest, McKinsey

**Rick D. Scruggs**, Chief Commercial Officer, Head of US Operations  
Salix, Watson, Oclassen

**June S. Almenoff**, MD, PhD, Chief Medical Officer  
Furiex, GSK, Tigenix

**Bob J. Gilkin**, Senior VP Market Access & Trade Relations  
Synergy Pharmaceuticals, AstraZeneca

**Reginald Williams**, VP Quality  
Salix, Amgen, Eli Lilly and Merck & Co

**Danielle Abramson**, PhD, Senior VP Global Head of IP  
Brown University, Greenberg Traurig

**Craig Miller**, VP Trade Channel  
Salix, Oclassen Pharmaceuticals, Watson Pharmaceuticals

**Michelle Snelling**, VP Human Resources  
Salix Pharmaceuticals, INC Research

**Ben Martie**, VP Legal Affairs  
ViiV Healthcare (GSK), Chimerix Inc.



# Board of Directors and Advisory Board

## Board of Directors

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**Dror Ben-Asher, CEO**

P.C.M.I. Ltd.

**Eric Swenden**

Alterphama nv, Lifeline Scientific

**Kenneth Reed, MD**

Dermatologist; Director Minerva Biotechnologies

**Alessandro Della Chà**

Cosmo Pharma, Acacia Pharma Group plc

**Rick D. Scruggs**

Salix, Watson, Oclassen

**Ofer Tsimchi**

Danbar, Polysack, Director in several companies

**Alla Felder**

Neuroderm, PwC

**Shmuel Cabilly, PhD**

Scientist, Director in several life-science companies

## Advisory Board

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**Jeff Leighton, PhD**

Glaxo, Exogen, Genesis,  
Inspire

**Werner Tschollar, MD**

Past SVP for Worldwide R&D,  
Novartis

**Scott Harris, MD**

Lyric, Avaxia, Ocera, Napo

**Prof. Ran Oren, MD**

Digestive and Liver expert,  
Hadassah

**Prof. Chezy Barenholz, PhD**

Prof., Hebrew University of  
Jerusalem, Co-inventor of Doxil®

**Abe Schwartz**

Covalon Tech., CEO Cedara  
Software

**Prof. Colin Blakemore, PhD**

Oxford, Past CEO - UK Medical  
Research Council

**Ira Kalfus, MD**

Lev pharma, Aetna/US  
Healthcare

**Prof. Thomas Borody, MD**

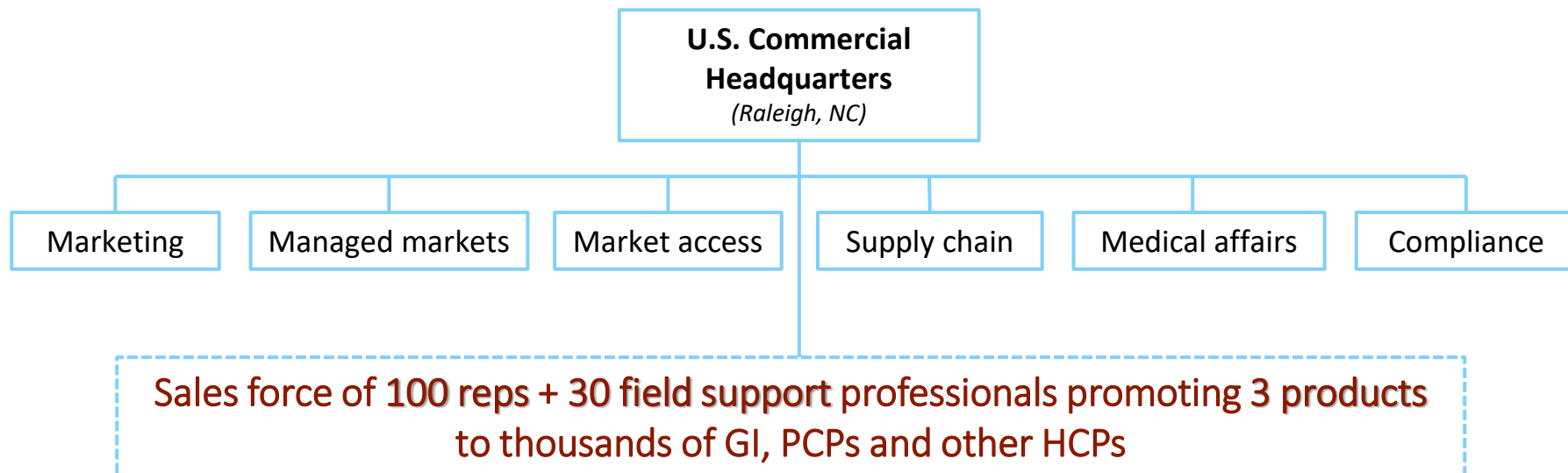
Founder Centre for Digestive Diseases

**Jerry Rosenblatt, PhD**

Foster Rosenblatt Consulting, IMS Health

## Commercial Operations - Strong U.S. Presence

Launched Aemcolo® in Q4/2019, Talicia® in Q1/20 and promoting Movantik® since April 2020



 **Talicia**<sup>®</sup>  
(omeprazole magnesium,  
amoxicillin, and rifabutin)  
delayed-release capsules

 **movantik**<sup>®</sup> <sup>i</sup>  
(naloxegol) 25 mg tablets

 **Aemcolo**<sup>®</sup>  
(rifamycin) delayed  
release tablets

<sup>i</sup> Movantik® is a registered trademark of AstraZeneca



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**Indicated for the treatment of opioid-induced constipation (OIC) in adults with chronic non-cancer pain**

# MOVANTIK® (naloxegol)

## Important Safety Information

### IMPORTANT SAFETY INFORMATION

- **MOVANTIK may cause serious side effects, including:**
  - Opioid withdrawal. You may have symptoms of opioid withdrawal during treatment with MOVANTIK, including sweating, chills, diarrhea, stomach pain, anxiety, irritability, and yawning. Patients taking methadone to treat their pain may be more likely to experience stomach pain and diarrhea. Tell your doctor if you have any of these symptoms
  - Severe Stomach Pain and/or Diarrhea. This can happen within a few days of starting MOVANTIK and can lead to hospitalization. If either of these side effects occurs, stop taking MOVANTIK and call your doctor immediately
  - Tear in your stomach or intestinal wall (perforation). Stomach pain that is severe can be a sign of a serious medical condition. If you get stomach pain that gets worse or does not go away, stop taking MOVANTIK and get emergency medical help right away
- **Do not take MOVANTIK if you:**
  - Have a bowel blockage (intestinal obstruction) or have a history of bowel blockage
  - Are allergic to MOVANTIK or any of the ingredients in MOVANTIK
- MOVANTIK can interact with other medicines and cause side effects, including opioid withdrawal symptoms (see symptoms above). Tell your doctor or pharmacist before you start or stop any medicines during treatment with MOVANTIK
- **Before you take MOVANTIK, tell your doctor about all of your medical conditions, including if you:**
  - Have any stomach, bowel (intestines) problems, including inflammation in parts of the large intestine (diverticulitis), or inflammation and injury of the intestines caused by reduced blood flow (ischemic colitis)
  - Have had recent surgery on the stomach or intestines
  - Have any kidney, or liver problems
  - Are pregnant or plan to become pregnant. Taking MOVANTIK during pregnancy may cause opioid withdrawal symptoms in you or your unborn baby. Tell your healthcare provider right away if you become pregnant during treatment with MOVANTIK
  - Are breastfeeding or plan to breastfeed. It is not known if MOVANTIK passes into your breast milk. Taking MOVANTIK while you are breastfeeding may cause opioid withdrawal in your baby. You and your healthcare provider should decide if you will take MOVANTIK or breastfeed. You should not breastfeed if you take MOVANTIK
- **Tell your doctor about all of the medicines you take**, including prescription and over-the-counter medicines, vitamins, and herbal supplements. Other medicines may affect the way MOVANTIK works
- **If you stop taking your opioid pain medicine, stop taking MOVANTIK and tell your doctor**
- Avoid eating grapefruit or drinking grapefruit juice during treatment with MOVANTIK
- **The most common side effects of MOVANTIK include:** Stomach (abdomen) pain, diarrhea, nausea, gas, vomiting, headache, and excessive sweating

### APPROVED USE FOR MOVANTIK

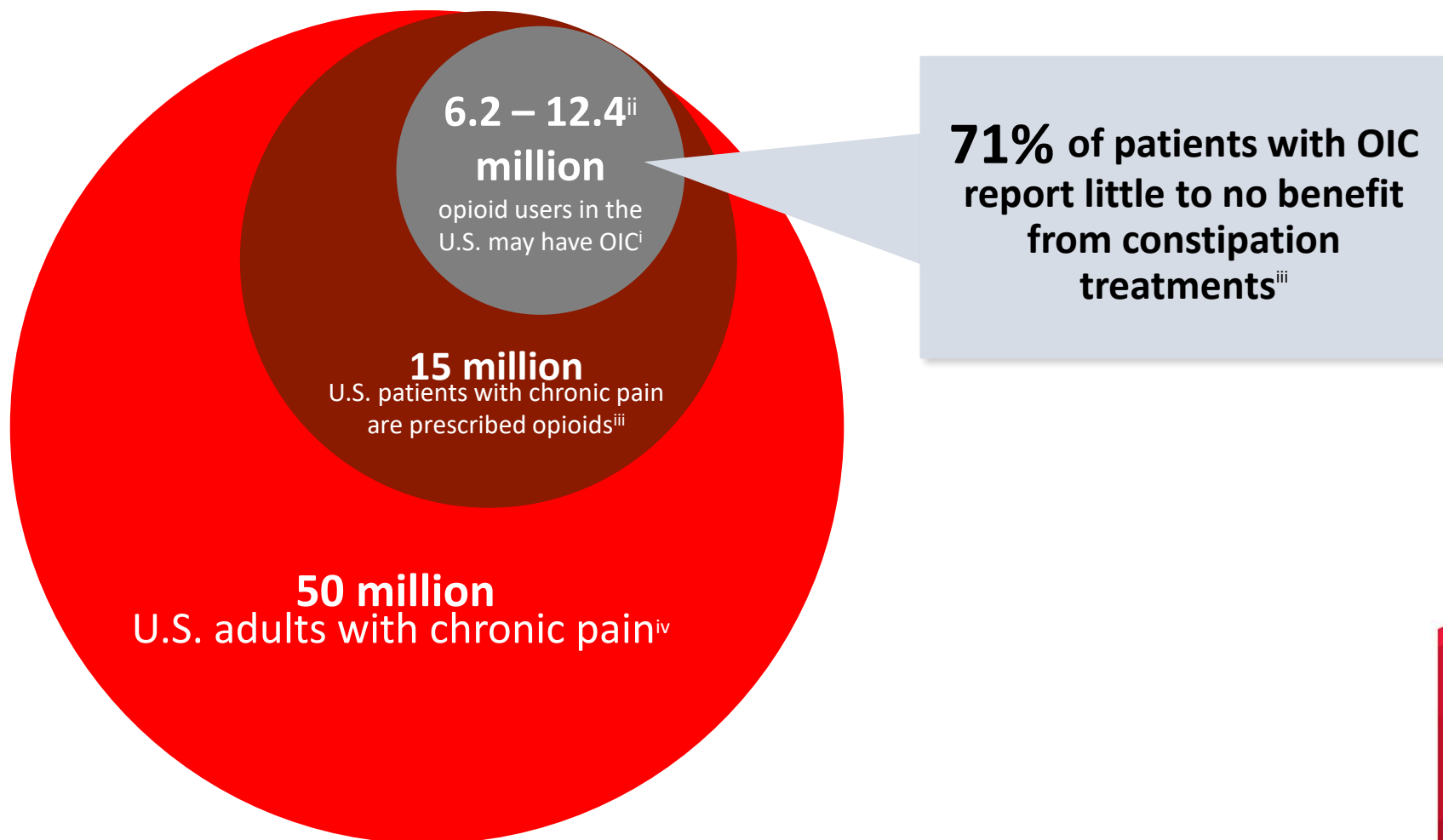
MOVANTIK is a prescription medicine used to treat constipation that is caused by prescription pain medicines called opioids, in adults with long-lasting (chronic) pain that is not caused by active cancer.

See full prescribing information for Movantik®: [www.movantik.com](http://www.movantik.com)

# Movantik® - FDA-Approved for Treatment of Opioid-Induced Constipation

<b>Approved Indication</b>	Treatment of opioid-induced constipation (OIC) in adults with chronic non-cancer pain
<b>Drug</b>	Oral naloxegol tablets available in 12.5mg and 25mg dosage strengths
<b>Approved</b>	Approved in the U.S. in 2014 - launched in 2015 by AstraZeneca and Daiichi Sankyo
<b>Key Attributes</b>	<ul style="list-style-type: none"> <li>✓ Specifically designed for opioid-induced constipation</li> <li>✓ Favorable tolerability and safety profile</li> <li>✓ Available in two doses</li> <li>✓ Strong reimbursement coverage</li> </ul>
<b>Market Exclusivity</b>	Patent protection extending until at least 2028

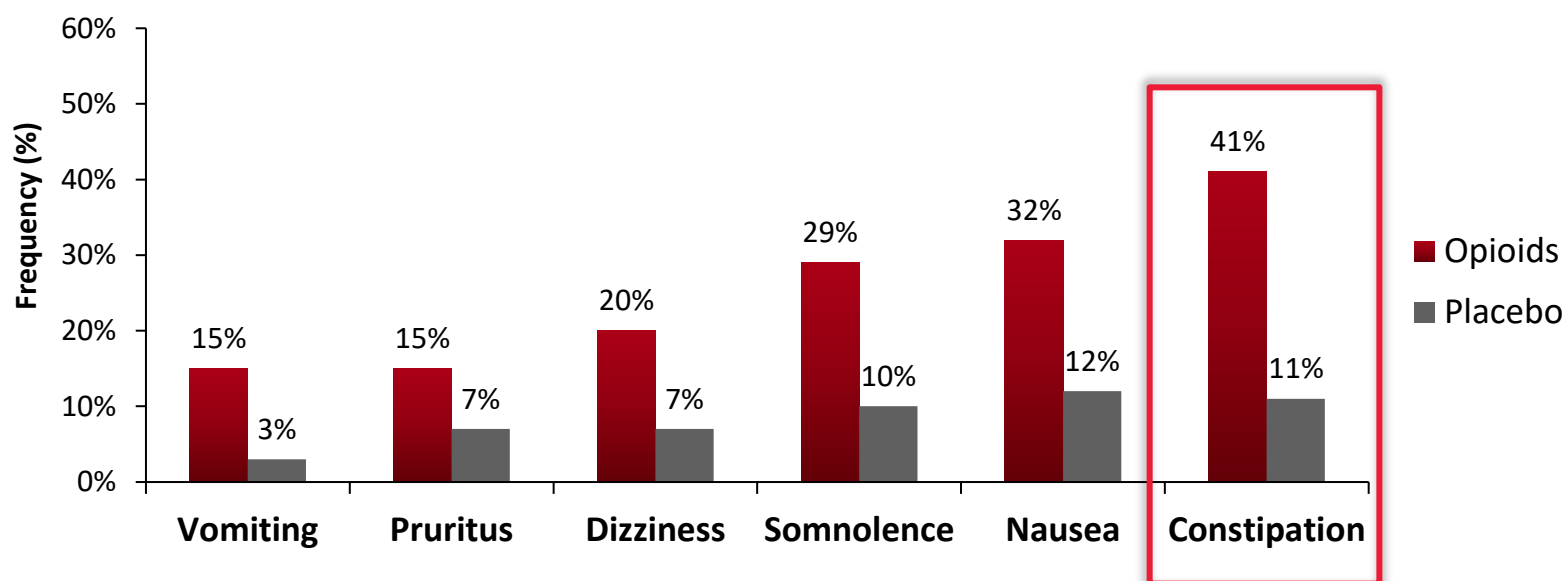
# OIC is a Large and Underserved Market<sup>i</sup>



<sup>i</sup> Bell TJ et al. *Pain Med.* 2009;10(1):35-42. Dahlhamer J, Lucas J, Zelaya C et al. Prevalence of Chronic Pain and High-Impact Chronic Pain Among Adults – United States, 2016. *MMWR Morb Mortal Wkly Rep.* 2018;67:1001-1006. Accessed September 29, 2020. <https://www.cdc.gov/mmwr/volumes/67/wr/pdfs/mm6736a2-H.pdf>. <sup>ii</sup> 30.7% of patients with chronic noncancer pain are prescribed opioids. Prevalence of OIC is estimated at between 40% to 81% in patients with chronic noncancer pain. Mathieson S et al. *J Int Med.* 2020;287:458-474. <sup>iii</sup> Constipation treatments included OTC laxatives (stool softeners, osmotics, stimulants, salines, and rectal options), prescription laxatives, and behavioral therapies (fiber supplements, increased fluids and exercise, and dietary changes). Coyne KS et al. *Clinicoecon Outcomes Res.* 2014;6:269-281. <sup>iv</sup> Vegia AR et al. *Pain Res Treat.* 2018. doi: 10.1155/2018/5704627 5. Coyne KS et al. *Clinicoecon Outcomes Res.* 2014;6:269-281.

# Constipation is One of the Most Common Side Effects of Opioid Therapy<sup>i</sup>

Results from a meta-analysis that included randomized controlled trials of opioid therapy in patients with chronic non-cancer pain:<sup>ii</sup>



***OIC Can Occur with Initiation of Opioid Therapy and May Persist for the Duration of Treatment<sup>j</sup>***

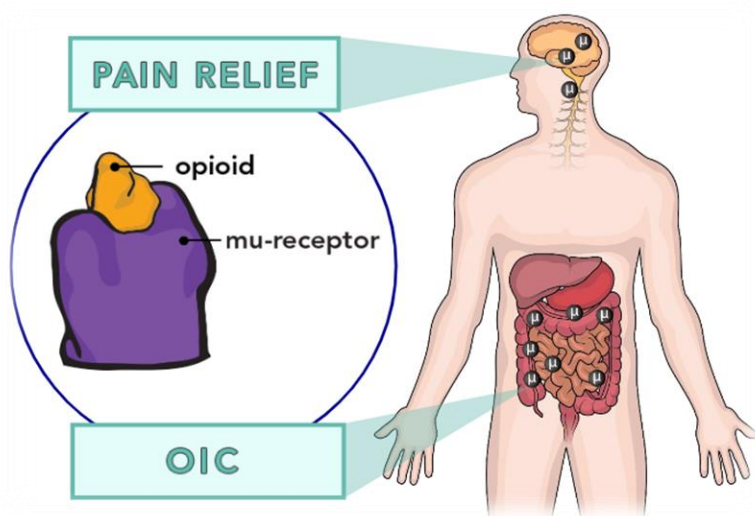
<sup>i</sup> Panchal SJ et al. *Int J Clin Pract.* 2007;61:1181-1187. Becker G et al. *Lancet.* 2009;373:1198-1206.

<sup>ii</sup> Kalso E et al. *Pain.* 2004;112:372-380.

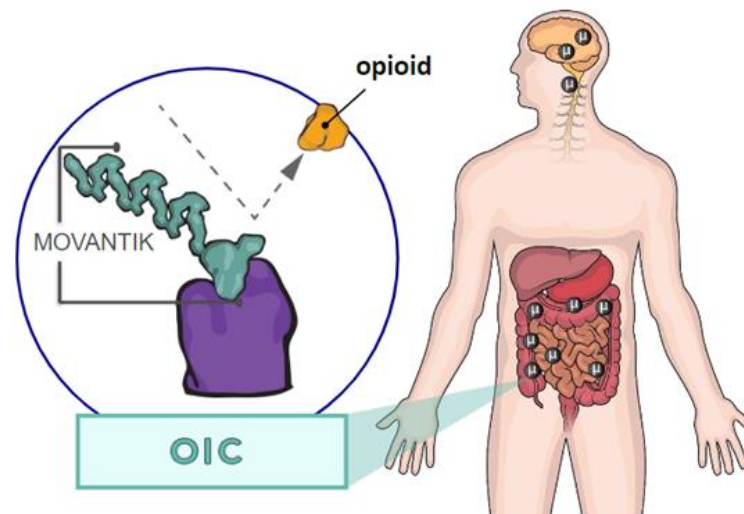
# Movantik® is a Once-Daily Oral Therapy Specifically Designed to Treat OIC

Movantik functions as a peripherally acting mu-opioid receptor antagonist (PAMORA) in tissues such as the bowel, thereby decreasing the constipating effect of opioids while limiting the potential for interference with centrally mediated opioid analgesia.

*OIC is caused by opioid binding to mu-receptors in the GI tract*



*Movantik antagonizes opioid binding at the mu-receptor\**



*Movantik is a PEGylated derivative of naloxone, limiting central effect<sup>i</sup>*



# Movantik - the #1 Prescribed Oral PAMORA<sup>i</sup>



Over **2,000,000** prescriptions written since 2015<sup>ii</sup>



The **#1 prescribed** oral PAMORA specifically designed to treat OIC<sup>ii</sup>



The American Gastroenterological Association recommends the use of Movantik as one of the prescription options for management of OIC<sup>iii</sup>



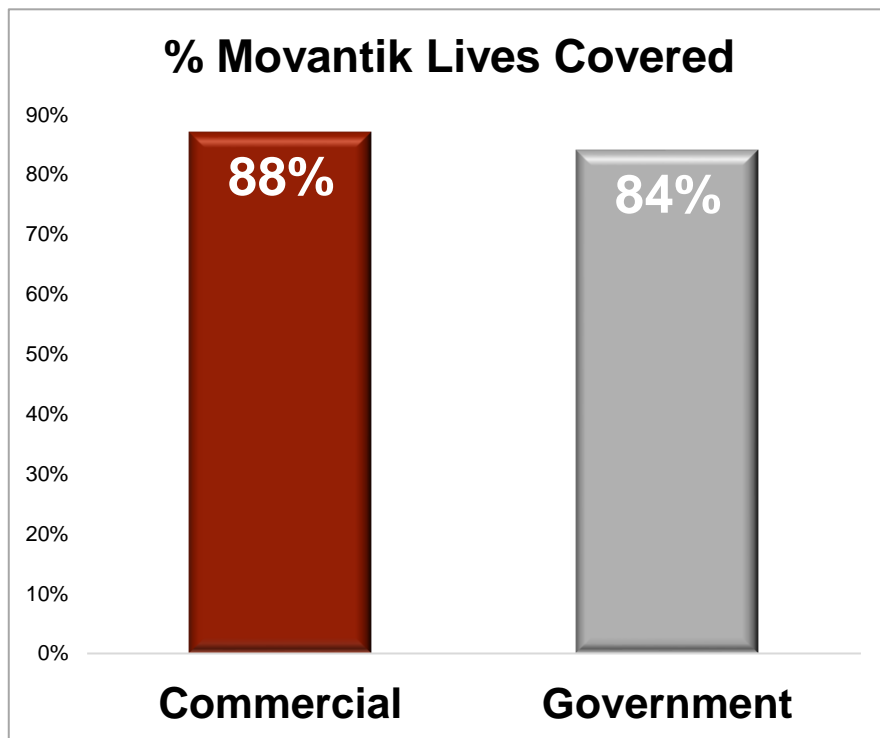
Movantik is covered or preferred without prior authorization for the majority of commercial and Medicare Part D patients in the U.S.<sup>iv</sup>

# Movantik Acquisition - A Transformative Event for RedHill

**RedHill acquired the global rights to Movantik<sup>i</sup> (excluding Europe and Canada) from AstraZeneca in April 2020**

- RedHill benefits from the large investment made by AstraZeneca to make Movantik a brand leader
  - First oral PAMORA approved in the U.S. for the treatment of OIC
- **RedHill is enhancing focus to grow this product**
  - Three consecutive quarters of Movantik prescription (TRx) growth led by RedHill promotion, reversing the trend of prescription decline prior to RedHill acquisition
  - RedHill's sales force is enlarging promotional footprint
  - Targeting gastroenterologists, primary care physicians and additional specialists

## Continued Best Coverage in PAMORA Class



- ✓ Best coverage without restrictions in the PAMORA class for both Commercial and Government segments
- ✓ 9 out of 10 Commercially insured patients; and 7 out of 10 Medicare Part D patients can access Movantik
- ✓ We have identified coverage opportunities and anticipate growth in 2021 for both segments

# **Talicia**®

(omeprazole magnesium,  
amoxicillin, and rifabutin)  
delayed-release capsules

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**Indicated for the treatment of *Helicobacter pylori* infection in adults**

***Approved for Marketing by U.S. FDA Following Two Positive Phase 3 Studies  
Launched in the U.S.***



## **Talicia® (omeprazole magnesium, amoxicillin and rifabutin)**

### **Important Safety Information**

Talicia contains omeprazole, a proton pump inhibitor (PPI), amoxicillin, a penicillin-class antibacterial, and rifabutin, a rifamycin antibacterial. It is contraindicated in patients with known hypersensitivity to any of these medications, any other components of the formulation, any other beta-lactams or any other rifamycins.

Talicia is contraindicated in patients receiving delavirdine, voriconazole or rilpivirine-containing products.

Serious and occasionally fatal hypersensitivity reactions have been reported with omeprazole, amoxicillin and rifabutin.

Acute Tubulointerstitial Nephritis has been observed in patients taking PPIs and penicillins.

Clostridioides difficile-associated diarrhea has been reported with use of nearly all antibacterial agents and may range from mild diarrhea to fatal colitis.

Talicia may cause fetal harm and is not recommended for use in pregnancy. It may also reduce the efficacy of hormonal contraceptives. An additional non-hormonal method of contraception is recommended when taking Talicia.

Talicia should not be used in patients with hepatic impairment or severe renal impairment.

Cutaneous lupus erythematosus and systemic lupus erythematosus have been reported in patients taking PPIs. These events have occurred as both new onset and exacerbation of existing autoimmune disease.

The most common adverse reactions ( $\geq 1\%$ ) were diarrhea, headache, nausea, abdominal pain, chromaturia, rash, dyspepsia, oropharyngeal pain, vomiting, and vulvovaginal candidiasis.

# Talicia® - Approved by U.S. FDA for Treatment of *H. pylori* Infection in Adults

Approved Indication	Treatment of <i>H. pylori</i> infection in adults
Drug	Omeprazole magnesium, amoxicillin and rifabutin 10 mg <sup>i</sup> / 250 mg/ 12.5 mg delayed-release capsules <sup>i</sup>
Approved	<b>NDA approved by U.S. FDA in November 2019 - launched in March 2020</b>
Key Attributes	<ul style="list-style-type: none"> <li>✓ Addresses concerns of resistance to clarithromycin and metronidazole</li> <li>✓ Favorable tolerability and safety profile</li> <li>✓ Aims to become the new first-line standard-of-care with broader indication</li> <li>✓ All-in-one capsule: supports ease of adherence and compliance, with a single co-pay</li> </ul>
Market Exclusivity	<ul style="list-style-type: none"> <li>– <b>Eligible for extended market exclusivity for total of 8 years</b> under QIDP designation</li> <li>– Patent protection extending until at least 2034</li> </ul>
Market Size	<ul style="list-style-type: none"> <li>– <b>Affects over 50% of the world population, with ~2 million U.S. patients treated annually</b></li> <li>– 2018 U.S. and global markets estimated at up to \$1.4 billion and \$4.8 billion respectively<sup>ii</sup></li> <li>– <b>Approximately 300% quarter-over-quarter prescription growth</b> and rapid expansion of the prescriber base</li> <li>– <b>National coverage for 167 million lives</b>, with additional coverage expected</li> </ul>

<sup>i</sup> Each delayed-release capsule contains omeprazole 10 mg (equivalent to 10.3 mg omeprazole magnesium), amoxicillin 250 mg, and rifabutin 12.5 mg

<sup>ii</sup> Foster Rosenblatt market analysis, October 2018

## Talicia Field Promotion Initiated July 2020

RedHill's U.S. sales force promoting Talicia® to approx. 25,000 gastroenterologists,  
primary care physicians and other healthcare providers  
Rapid expansion in managed care coverage since launch

**Talicia®**  
(omeprazole magnesium,  
amoxicillin, and rifabutin)  
delayed-release capsules

Full Prescribing Information

Important Safety Information

Talicia contains omeprazole, a proton pump inhibitor (PPI),  
amoxicillin a penicillin-class antibacterial and rifabutin, a

[Home](#) [The Challenge of \*H. pylori\*](#) [Efficacy](#) [Safety](#) [Dosing & Administration](#) [Information for Patients](#) [Savings Program](#)

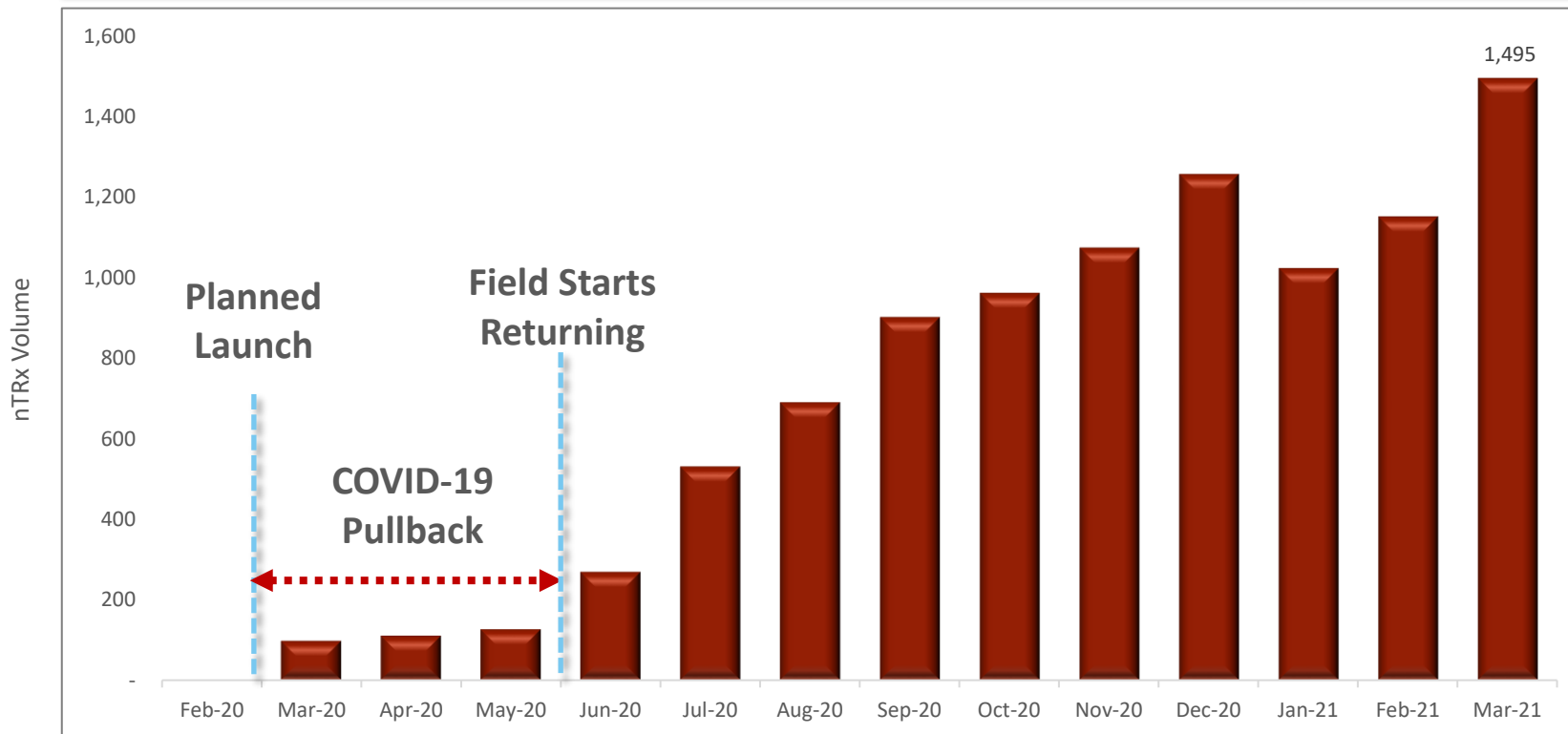
For the treatment of *Helicobacter pylori* infection in adults

**Outsmart Resistance.  
Eradicate *H. pylori*.**



# Talicia: Strong nTRx Volume Growth With New High for nTRx Volume in March

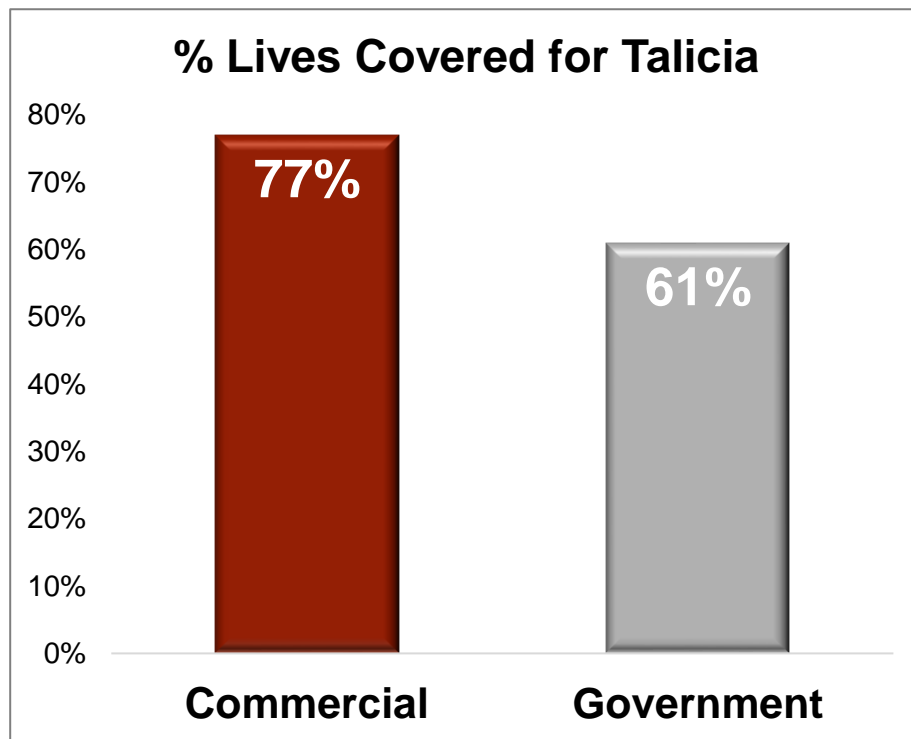
## Talicia nTRx Growth



**Continued to achieve new launch year milestones; 11% prescription growth in Q1/21 and anticipating accelerated brand growth in coming months**



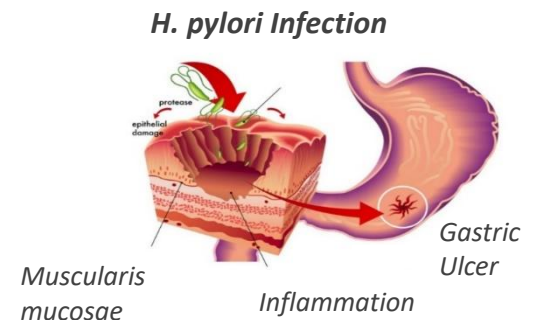
## Talicia Commercial Coverage Reached 69%



- ✓ Talicia Commercial coverage continues to grow - coverage has reached 77% (May 2021)
- ✓ 8 out of 10 Commercially insured patients can access Talicia, with most having access without restrictions
- ✓ Payer strategy continues to support expanding coverage

## *H. pylori* infection → Progressive gastro-duodenal damage

- *H. pylori* - a Group I carcinogen and the strongest risk factor for gastric cancer<sup>i</sup> and peptic ulcer disease; Also associated with iron deficiency, B12 deficiency and drug malabsorption
  - Gastric cancer - a leading cause of cancer mortality worldwide, accounting for ~700K deaths annually
  - Eradication of *H. pylori* infection reduces gastric cancer risk<sup>ii</sup> by about 75%
- U.S. *H. pylori* prevalence estimated at approx. 35% of the population - over 100 million people<sup>iii</sup>, with an estimated 2 million patients treated annually<sup>iv</sup>
- SoC fails in approximately 25-40% of patients due to growing resistance to clarithromycin and metronidazole - antibiotics commonly used in standard combination therapies<sup>v</sup>
- **American College of Gastroenterology (ACG) guidelines generally exclude the majority of U.S. population from treatment with SoC** - recommending against clarithromycin-based triple SoC therapies in cases of prior macrolide exposure (most of the U.S. population), in regions with unknown clarithromycin resistance (most U.S. regions) or regions with  $\geq 15\%$  clarithromycin resistance<sup>vi</sup>
  - Failed antibiotic treatments create newly-resistant bacteria<sup>vii</sup>



<sup>i</sup> Lamb A et al. Role of the *Helicobacter pylori*-induced inflammatory response in the development of gastric cancer. *J Cell Biochem* 2013 Mar;114(3):491-7; <sup>ii</sup> Kumar S et al. Risk Factors and Incidence of Gastric Cancer After Detection of *Helicobacter pylori* Infection: A Large Cohort Study, *Gastroenterology* 2019. <sup>iii</sup> Hooi JKY et al. Global Prevalence of *Helicobacter pylori* Infection: Systematic Review and Meta-Analysis. *Gastroenterology* 2017;153:420-429; <sup>iv</sup> IQVIA Custom Study for RedHill Biopharma, 2019; <sup>v</sup> Malfertheiner P. et al. Management of *Helicobacter pylori* infection - the Maastricht IV/ Florence Consensus Report, *Gut* 2012;61:646-664; O'Connor A. et al. Treatment of *Helicobacter pylori* Infection 2015, *Helicobacter* 20 (S1) 54-61; Venerito M. et al. Meta-analysis of bismuth quadruple therapy versus clarithromycin triple therapy for empiric primary treatment of *Helicobacter pylori* infection. *Digestion* 2013;88(1):33-4; <sup>vi</sup> Chey, WD et al. *Am. J. Gastroenterol* 2017; 112: 212-38. <sup>vii</sup> Nishizawa T et al. Enhancement of Amoxicillin Resistance after Unsuccessful *Helicobacter pylori* Eradication. *Antimicrob. Agents Chemother.* 2011, p. 3012-3014.

# *H. pylori* Resistance - An Important Public Health Concern

## WHO: *H. pylori* ranked as pathogen with “high priority” need for new treatments

- February 2017 - WHO publishes global priority pathogen list
- Intended to identify the most important resistant bacteria at a global level for which there is an urgent need for new treatments
- H. pylori* (clarithromycin-resistant) categorized as a pathogen for which there is a High Priority need to develop new treatments

*Enterococcus faecium*,  
vancomycin-resistant  
*Staphylococcus aureus*,  
methicillin-resistant,  
vancomycin intermediate  
and resistant

*Helicobacter pylori*,  
clarithromycin-resistant

*Campylobacter*,  
fluoroquinolone-resistant

*Salmonella spp.*,  
fluoroquinolone-resistant

*Neisseria gonorrhoeae*,  
3rd generation  
cephalosporin-resistant,  
fluoroquinolone-resistant

## FDA: *H. pylori* identified under the GAIN Act as pathogen posing serious threat to public health

- Talicia® received FDA QIDP designation under the GAIN Act for serious or life-threatening infections
- Priority Review
- Extended market exclusivity for a total of 8 years

The three new qualifying pathogens are:

*Coccidioides species*

*Cryptococcus species*

*Helicobacter pylori*

All 18 of the original draft pathogens remain on the list

# Positive Pivotal Phase 3 Study

Randomized, Double-Blind, Active Comparator, Two-Arm, Pivotal Phase 3 Study (ERADICATE Hp2), Comparing Talicia® Against a Regimen of Amoxicillin and Omeprazole Alone in the Treatment of Confirmed *H. pylori* Infection in Dyspepsia Patients, Regardless of Ulcer Status

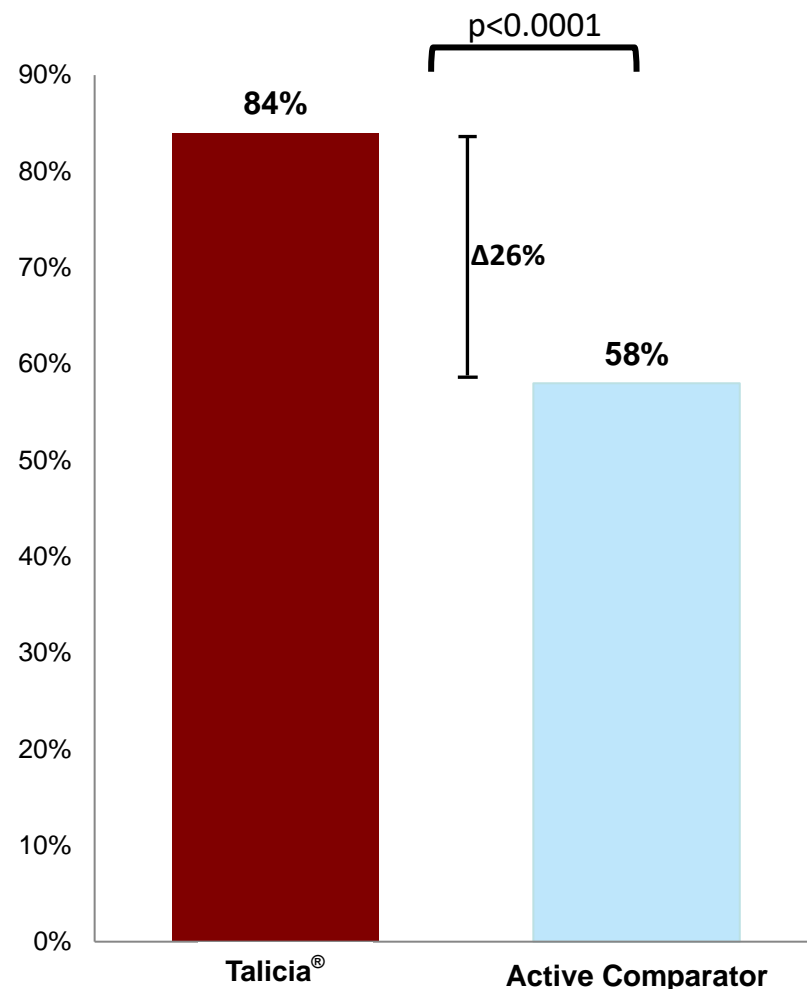
<b>Number of subjects</b>	455 in the U.S.
<b>Duration of Study Treatment</b>	14 days
<b>Primary Endpoint</b>	- The occurrence of <i>H. pylori</i> eradication in the Talicia® group compared to the active comparator group as confirmed via <sup>13</sup> C Urea Breath Test (UBT) testing 43-71 days after initiation of treatment

# Positive Pivotal Phase 3 Study

The ERADICATE Hp2 study met its primary endpoint with a high degree of statistical significance ( $p < 0.0001$ )

- Top-line results demonstrated 84% eradication of *H. pylori* with Talicia® vs. 58% in the active comparator arm ( $p < 0.0001$ ) (ITT)
- **90% eradication** based on adherence analysis (protocol defined subpopulation with evidence of drug exposure: designated as PK population)
- No serious safety issues were reported in the study; Talicia® found to be well-tolerated

Analysis*	Phase 3 Results - Talicia vs. Active Comparator	p-value
ITT Population	84% vs. 58%	$p < 0.0001$
mITT Population	84% vs. 58%	$p < 0.0001$
<b>PK population</b> (evidence of drug exposure)	<b>90% vs. 65%</b>	<b><math>P &lt; 0.0001</math></b>



\* ITT population included all randomized patients who received at least one dose of study drug; PK population included those subjects in the ITT population who had demonstrated presence of any component of investigational drug at Visit 3 (approx. day 13) or had undetected levels drawn >250 hours after the last dose.

# Resistance Patterns in Pivotal Study Demonstrate Zero to Negligible Resistance For Component Antibiotics in Talicia®

## *H. pylori* Resistance to Standard-of-Care

- H. pylori* culture results from patients across 20 U.S. states supported the high resistance of *H. pylori* to the antibiotics most commonly used for treatment -

Antibiotic	<i>H. pylori</i> Resistance Rate
Metronidazole	44%
Clarithromycin	17%
Amoxicillin	6%
<b>Rifabutin</b>	<b>0%</b>

- Consistent with the literature<sup>i</sup> describing the diminished efficacy of standard-of-care therapies, open-label part of the study showed 53% eradication of *H. pylori* with standard-of-care<sup>ii</sup>
- Consistent 21-29% treatment benefit of Talicia® vs. the active comparator across all *H. pylori* culture susceptibility and resistance subgroups

<sup>i</sup> Fallone CA et al. The Toronto Consensus for the Treatment of Helicobacter pylori Infection in Adults. Gastroenterology 2016;151:51–69.

<sup>ii</sup> N=90; therapy in open-label part with patients who failed eradication was determined by the treating physician

## Positive First Phase 3 Study

A Randomized Placebo-Controlled Phase 3 Study (ERADICATE Hp) to Assess the Safety and Efficacy of Talicia® in the Treatment of Confirmed *H. pylori* Infection in Dyspepsia Patients, Regardless of Ulcer Status

<b>Number of Subjects</b>	118 in the U.S.
<b>Treatment Duration</b>	14 days
<b>Primary Endpoints</b>	<ul style="list-style-type: none"> <li>- The occurrence of <i>H. pylori</i> eradication as confirmed via 13C UBT testing 28-35 days after completion of treatment</li> <li>- Superiority in eradication of <i>H. pylori</i> infection over historical standard of care efficacy levels of 70% effectiveness</li> </ul>
<b>Positive Phase 3 Results</b>	<ul style="list-style-type: none"> <li>- <b>Study met protocol-defined primary endpoint - demonstrating 89.4% efficacy in eradicating <i>H. pylori</i> infection (<math>p &lt; 0.001</math>)</b></li> <li>- <b>63% eradication rate demonstrated in open-label treatment of treatment naïve placebo-arm patients with physician choice therapy (mostly standard of care (SoC)) for persistent <i>H. pylori</i> infection</b> - supporting the potential superior efficacy of Talicia</li> <li>- No serious adverse events related to the drug were noted in the study</li> </ul>

# Strong Protection from Competition and Substitution

- FDA QIDP designation providing eligibility for a total of **8 years of market exclusivity**
- Talicia is covered by U.S. patents covering the unique 'all-in-one' capsule formulation and distinct PK profile, **extending until at least 2034**, with additional pending patents and applications in various territories worldwide
- Clear dosing differentiation versus available commercial options; rifabutin dosing (50mg tid) not commercially available and cannot be replicated by prescribers
- Acute use with low sensitivity for price differences → low incentive for substitution
- Indication allowing unique labeling, marketing and promotional opportunity
- 'All-in-one' easy to use regimen with single co-pay preferred over more cumbersome regimens



# Talicia® - Clear Clinical Differentiation Provides Large Potential for Market Opportunity in the U.S. and WW

## High Prevalence of *H. pylori*

Over 27M treatments  
annually WW<sup>i</sup>:  
**U.S.:** 2M  
**5EU:** up to 3.2M  
**Japan:** up to 1.4M  
**China:** up to 4.1M

## Diminished Efficacy of Standard-of-Care - Approx. 60%

Growing *H. pylori*  
resistance has led to  
diminished efficacy of  
current standard-of-care

## Current Brand Medications Lack Clinical Differentiation

Current brands provide  
only modest convenience  
improvement vs. generics

## \$4.8B Global Market

Annual U.S. market for  
*H. pylori* therapies  
estimated at \$1.4B<sup>i</sup>

## Talicia® - Potential First-Line Therapy Targeting up to \$1.4B U.S. Market

- **Efficacy** - demonstrated clinical activity with high statistical significance in eradicating *H. pylori* in U.S. pivotal Phase 3 study
- **Addresses concerns of resistance** to clarithromycin and metronidazole
- **Attractive tolerability profile**
- **Potential to become preferred first-line treatment**
- **First all-in-one fixed-dose** - simple regimen potentially improves compliance and efficacy; Additional protection against generic substitution; Single co-pay



**Indicated for the treatment of travelers' diarrhea caused by noninvasive strains of *E. coli* in adults**

***Approved for Marketing by U.S. FDA  
Launched in the U.S. by RedHill - December 2019***



# Aemcolo® (rifamycin) - Important Safety Information

## INDICATION AND IMPORTANT SAFETY INFORMATION

Aemcolo® is indicated for the treatment of travelers' diarrhea caused by noninvasive strains of *Escherichia coli* (*E. coli*) in adults.

### Limitations of Use

Aemcolo® is not indicated in patients with diarrhea complicated by fever or bloody stool or due to pathogens other than noninvasive strains of *E. coli*.

To reduce the development of drug-resistant bacteria and maintain the effectiveness of Aemcolo® and other antibacterial drugs, Aemcolo® should be used only to treat or prevent infections that are proven or strongly suspected to be caused by susceptible bacteria.

## CONTRAINDICATION

Aemcolo® is contraindicated in patients with a known hypersensitivity to rifamycin, any of the other rifamycin class antimicrobial agents, or any of the components in Aemcolo®.

## WARNINGS AND PRECAUTIONS

**Risk of Persistent or Worsening Diarrhea Complicated by Fever and/or Bloody Stool**

Aemcolo® was not shown to be effective in patients with diarrhea complicated by fever and/or bloody stool or diarrhea caused by pathogens other than *E. coli* and is not recommended for use in such patients.

Discontinue Aemcolo® if diarrhea gets worse or persists more than 48 hours and consider alternative antibacterial therapy.

### Clostridium difficile-Associated Diarrhea (CDAD)

CDAD has been reported with use of nearly all antibacterial agents and may range in severity from mild diarrhea to fatal colitis.

Consider CDAD in all patients who present with diarrhea following antibacterial drug use. Careful medical history is necessary since CDAD has been reported to occur over two months after the administration of antibacterial agents.

### Development of Drug-Resistant Bacteria

Prescribing Aemcolo® in the absence of a proven or strongly suspected bacterial infection or a prophylactic indication is unlikely to provide benefit to the patient and increases the risk of the development of drug-resistant bacteria.

## ADVERSE REACTIONS

Discontinuation of Aemcolo® due to adverse reactions occurred in 1% of patients. The most frequent adverse reactions were abdominal pain (0.5%) and pyrexia (0.3%).

Adverse reactions that occurred in at least 2% of Aemcolo®-treated patients and with a higher incidence than in the placebo or ciprofloxacin groups were constipation 3.5% and headache 3.3%, respectively.

## USE IN SPECIFIC POPULATIONS

### Pregnancy

There are no available data on AEMCOLO use in pregnant women to inform any drug associated risks for major birth defects, miscarriage, or adverse maternal or fetal outcomes.

### Lactation

There is no information regarding the presence of AEMCOLO in human milk, the effects on the breastfed infant, or the effects on milk production.

### Pediatric Use

The safety and effectiveness of AEMCOLO has not been established in pediatric patients <18 years of age.

See Full prescribing information for Aemcolo® is available at [www.aemcolo.com](http://www.aemcolo.com)

## Launched by RedHill in the U.S. - December 2019

- A rifamycin antibacterial approved by U.S. FDA in Nov. 2018 for the treatment of travelers' diarrhea caused by noninvasive strains of *E. coli* in adults
- In-licensed U.S. rights from Cosmo Pharmaceuticals N.V. in Oct. 2019
- Robust U.S. patent portfolio and FDA QIDP designation, with U.S. marketing exclusivity through 2028

- ✓ Minimally absorbed
- ✓ Targeted delivery system
- ✓ Proven efficacy against *E. coli*
- ✓ Reliable safety and tolerability
- ✓ Simple BID dosing



# High Travel Spending Per Person in the U.S. Points to Significant Commercial Opportunity

## Travelers' diarrhea is a significant market:

- Approximately **93 million** Americans travel abroad in 2018, of which **60 million** traveled to medium to high risk regions<sup>i</sup>
- Travelers' diarrhea may affect up to **70%** of travelers depending on destination and season of travel<sup>ii</sup>
- **> 1/3** of those travelers are seeking health advice prior to leaving<sup>iii</sup>
- **52%** of travelers travel with OTC meds for gastro ailments and **28%** travel with a prescription antibiotic<sup>iv</sup>
- The International Society of Travel Medicine (ISTM) recommends **traveling with an antibiotic for self-treatment** when visiting developing regions<sup>iii</sup>

<sup>i</sup> <https://travel.trade.gov/view/m-2018-O-001/index.html>; <sup>ii</sup> CDC Yellow Book; <sup>iii</sup> "Travel Health, Knowledge, Attitudes and Practices among United States Travelers." J Travel Med 2004; 11:23–26; <sup>iv</sup> Aemcolo survey market research, Aries Sep. 2019;



## **Opaganib (ABC294640)<sup>i</sup>**

*Investigational new drug*

*First-in-class, orally-administered sphingosine kinase-2 (SK2) inhibitor  
targeting multiple oncology, inflammatory and GI indications*

***Enrollment of global Phase 2/3 study completed and positive top-line  
data from U.S. Phase 2 study for COVID-19***

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***Phase 2 study for the treatment of cholangiocarcinoma ongoing***

<sup>i</sup> Yeliva® is the proposed tradename for the drug product containing opaganib, which is subject to review by the FDA at the time of NDA filing

# Opaganib - SK2 Inhibitor for COVID-19 and Oncology, Gastrointestinal and Inflammatory Diseases

<b>The Product</b>	<ul style="list-style-type: none"> <li>- Potential first-in-class, orally-administered sphingosine kinase-2 (SK2) inhibitor - with anti-cancer, anti-inflammatory activities, targeting multiple oncology, inflammatory and GI indications</li> <li>- <b>Potent anti-viral activity, targeting a critical host factor - minimizing potential development of resistance due to viral mutations</b></li> </ul>
<b>Potential Market</b>	<ul style="list-style-type: none"> <li>- Significant market potential - multiple indications with an unmet need</li> </ul>
<b>Development Status</b>	<ul style="list-style-type: none"> <li>- Completed numerous successful pre-clinical studies in oncology, GI-Inflammation and radioprotection models, as well as food effect and toxicology studies</li> <li>- Phase 1 study in cancer patients with advanced solid tumors successfully met primary and secondary endpoints</li> <li>- <b>Phase 2a study for treatment of cholangiocarcinoma ongoing</b> <ul style="list-style-type: none"> <li>- Orphan Drug Designation for the treatment of cholangiocarcinoma</li> <li>- Compassionate use for cholangiocarcinoma under Expanded Access Program</li> </ul> </li> <li>- Investigator-sponsored Phase 2 study in prostate cancer initiated March 2020 at Medical University of South Carolina (MUSC) - supported by NCI grant to MUSC</li> <li>- <b>Enrollment, treatment and follow-up completed for the global Phase 2/3 study in patients hospitalized with severe COVID-19 pneumonia</b></li> <li>- <b>Reported positive top-line safety and efficacy data from U.S. Phase 2 study in patients hospitalized with COVID-19 pneumonia</b></li> </ul>

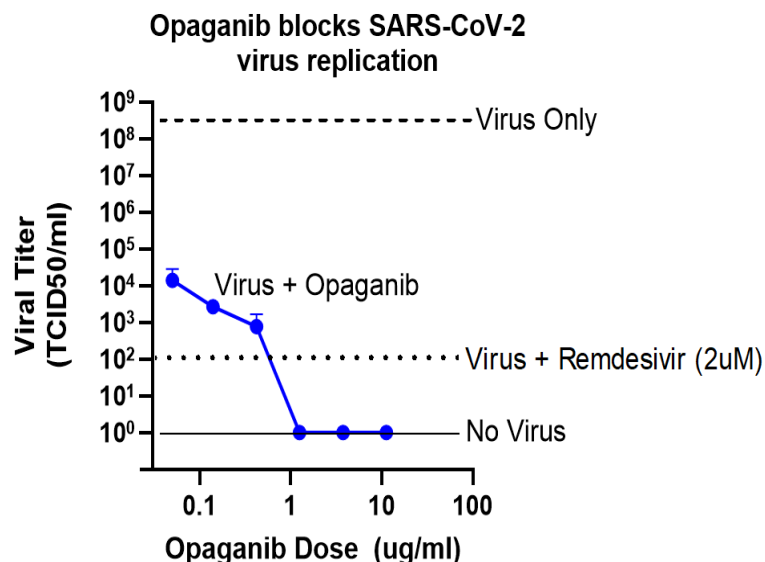
# Opaganib - Complete Inhibition of SARS-CoV-2

**Opaganib's unique MoA combines potent antiviral and anti-inflammatory activities, targeting a host cell component and minimizing likelihood of resistance**

## Potent Anti-SARS-CoV-2 Activity:

Opaganib evaluated in *in vitro* model of human lung bronchial tissue (EpiAirway™), including positive control of remdesivir:

- Opaganib completely inhibited SARS-CoV-2 viral replication as measured after three days incubation
- Opaganib demonstrated the most potent activity vs. all compounds tested, including remdesivir
- Dose-dependent inhibition of virus production without compromising cell membrane integrity
- Opaganib further demonstrated potent inhibition of Beta (South African) and Gamma (Brazilian) COVID-19 variants in preclinical study at non-cytotoxic doses





# Opaganib - COVID-19 Studies

## Global Phase 2/3 Study - Last Patient Out Announced July 19, 2021

Randomized, double-blind, parallel-arm, placebo-controlled global Phase 2/3 study

- 475 subjects enrolled with severe COVID-19 pneumonia in approx. 40 sites
- Primary endpoint: proportion of patients reaching room air by Day 14
- Study approved in 10 countries: Italy, UK, Russia, Israel, Mexico, Colombia, Poland, Brazil, Peru and the U.S.
- Four DSMB recommendations to continue study following independent safety and futility reviews

## U.S. Phase 2 Study - Completed

U.S. randomized, double-blind, placebo-controlled Phase 2 study

- Positive top-line safety and efficacy data
- Small sample size of 40 hospitalized patients with COVID-19 pneumonia
- Focused on safety and initial efficacy signals; not powered for statistical significance

- Enrollment, treatment and follow-up for the Phase 2/3 study completed
- Collaborations with U.S., European and Canadian suppliers for Manufacturing ramp-up

# Opaganib Phase 2 COVID-19 Study - Positive Top-Line Data

## U.S. Phase 2 study of opaganib in patients hospitalized with COVID-19 pneumonia:

Non-powered study of 40 patients, randomized 1:1 to received opaganib or placebo on top of standard-of-care. Follow up for up to 42 days post treatment initiation

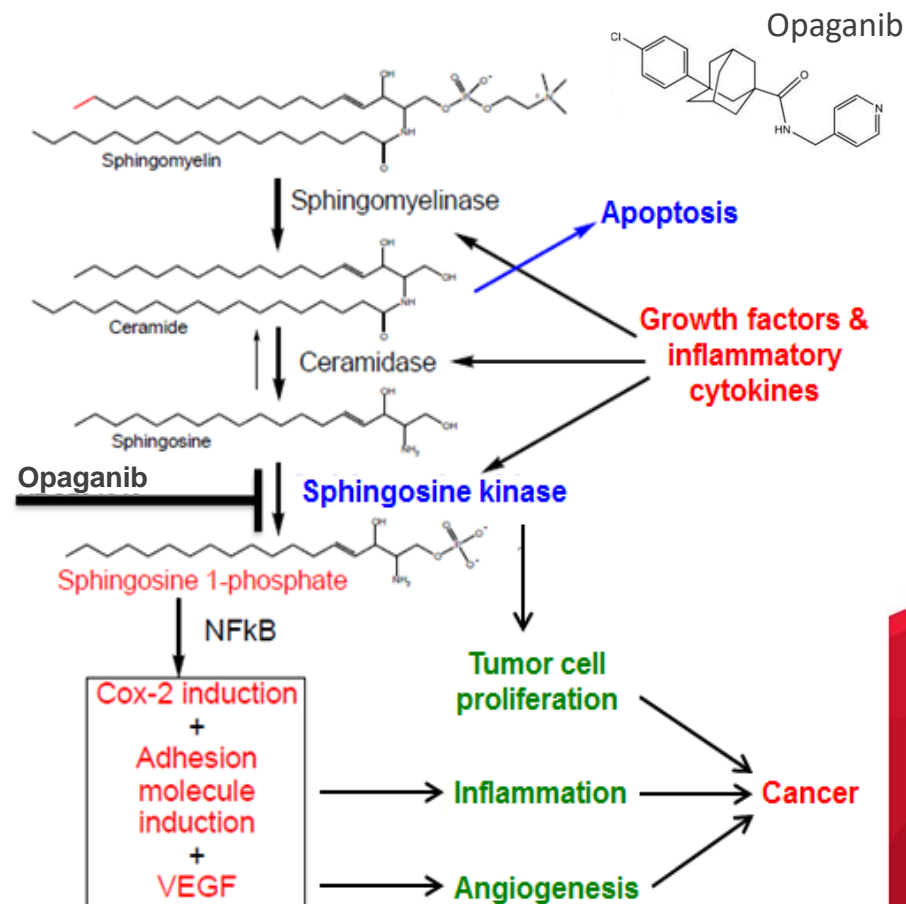
## Preliminary top-line data demonstrated positive safety and efficacy signals:

- Opaganib found to be safe, with no material safety differences between the arms. Overall, fewer patients suffered from SAEs in the opaganib arm vs. placebo
- Opaganib arm demonstrated consistent trend of greater improvement in reducing oxygen requirement by end of treatment on Day 14 across key primary and secondary efficacy outcomes:
  - ✓ 50% of opaganib treated patients reached room air by Day 14 vs. 22% in the placebo group – benefit maintained regardless of whether patients received dexamethasone and/or remdesivir
  - ✓ 86.4% of opaganib treated patients were discharged from hospital by Day 14 vs. 55.6% of patients treated with placebo
  - ✓ Median time to discharge was 6 days for the opaganib group compared to 7.5 days for the placebo group
  - ✓ 81.8% of opaganib patients achieved a 2-point improvement in the WHO Ordinal Scale vs. 55.6% of patients in the placebo group – achieved in a median time of 6 days vs. 7.5 days, respectively
  - ✓ Greater reduction from baseline of the median total oxygen requirement (AUC) over 14 days (62.0% vs. 47%)

# Opaganib - Novel SK2 Inhibitor with Anti-Cancer, Anti-Inflammatory and Anti-Viral Activities

**Opaganib selectively inhibits SK2, a lipid kinase with multiple cellular functions - potentially inhibiting tumor growth, pathological inflammation, and viral replication**

- Sphingosine kinase-2 (SK2) impacts tumor growth and proliferation, pathological inflammation and cytokine production, including TNF $\alpha$  and IL-6
- SK2 is a critical component of the replication-transcription complex of many viruses. Blocking it may play a role in inhibiting viral replication
- Opaganib demonstrated potent anti-viral activity, targeting a host cell component and minimizing likelihood of resistance



# Opaganib - Positive Phase 1 Study

## Phase 1 Clinical study of opaganib in Patients with Advanced Solid Tumors

<b>Study Design</b>	<ul style="list-style-type: none"> <li>- Open-label, dose-escalation, PK and PD, first-in-human Phase I study with opaganib</li> </ul>
<b>Site</b>	<ul style="list-style-type: none"> <li>- Medical University of South Carolina</li> </ul>
<b>Subjects</b>	<ul style="list-style-type: none"> <li>- 21 patients with advanced solid tumors, the majority of which were GI cancer patients</li> </ul>
<b>Endpoints</b>	<ul style="list-style-type: none"> <li>- Primary endpoints: to identify the maximum tolerated dose (MTD) and the dose limiting toxicities (DLTs) and to evaluate the safety of opaganib</li> <li>- The study included the first-ever longitudinal analyses of plasma S1P levels as a potential PD biomarker for activity of a sphingolipid-targeted drug</li> </ul>
<b>Positive Phase 1 Results (June 2016)</b>	<ul style="list-style-type: none"> <li>- The study met its primary and secondary endpoints</li> <li>- Opaganib was demonstrated to be safe and well tolerated and was shown to be safely administered to cancer patients at doses that provide circulating drug levels that are predicted to have therapeutic activity</li> <li>- Administration of opaganib resulted in a rapid and pronounced decrease in S1P levels</li> <li>- <b>One patient had a prolonged partial remission and several patients had prolonged stabilization of disease</b></li> <li>- Of the three patients with cholangiocarcinoma, one had a partial response and the other two stable disease, one which lasted for over a year</li> </ul>

# Opaganib - Phase 2a Study for Cholangiocarcinoma

## Ongoing

**Single-arm Phase 2a clinical study evaluating opaganib in patients with advanced, unresectable, intra-hepatic, perihilar and extra-hepatic cholangiocarcinoma**

<b>Sites</b>	Mayo Clinic, MD Anderson, Huntsman Cancer Institute, Emory University
<b>Initiated</b>	December 2017
<b>Lead Investigator</b>	Dr. Mitesh J. Borad, MD, Associate Professor of Medicine and Director of Phase I Drug Development at the Mayo Clinic Cancer Center in Arizona
<b>Development status</b>	<ul style="list-style-type: none"> <li>- Enrollment completed for stand-alone arm (39 subjects) - preliminary data indicates efficacy signal in a number of subjects; data to be submitted for presentation</li> <li>- Recruitment initiated for second arm evaluating opaganib in combination with hydroxychloroquine, an anti-autophagy agent</li> <li>- Addition of third arm planned, evaluating opaganib in combination with RHB-107, subject to discussions with FDA</li> </ul>
<b>Primary endpoint</b>	<ul style="list-style-type: none"> <li>- Response Rate - defined as CR+PR+SD of at least 4 months duration</li> </ul>
<b>Market Size</b>	<ul style="list-style-type: none"> <li>- Approximately 8,000 people are diagnosed with intrahepatic and extrahepatic bile duct cancers annually in the U.S.</li> </ul>



## **RHB-107 (upamostat)**

*Investigational new drug*

*Potential first-in-class small molecule targeting oncology, inflammatory lung diseases  
and GI indications*

***U.S. Phase 2/3 COVID-19 ongoing in non-hospitalized patients***

# RHB-107 - S1 Serine Protease Inhibitor with Ongoing Phase 2/3 COVID-19 Study in Non-Hospitalized Patients

The Drug	Potential first-in-class, orally-administered inhibitor of S1 family of trypsin-like serine proteases with potential for use in the treatment of cancer, inflammatory lung diseases, irritable bowel syndrome, inflammatory bowel disease and pancreatitis
	RHB-107 is a specific and potent inhibitor of human trypsin-3 ( $K_i \sim 20\text{nM}$ ), trypsin-2 ( $K_i \sim 75\text{nM}$ ), trypsin-6 ( $\sim 100\text{nM}$ ), trypsin-1 ( $K_i \sim 190\text{nM}$ ) and matriptase-1 ( $\sim 200\text{nM}$ )
	Licensed worldwide rights from Heidelberg Pharma (formerly Wilex), excluding China, Taiwan, Macao and Hong Kong
Development Status	<b>Demonstrated clinical safety profile from approx. 200 patients across 10 clinical studies, including Phase 2 studies in locally advanced pancreatic cancer and metastatic breast cancer</b>
	<ul style="list-style-type: none"> <li>– FDA Orphan Drug Designation awarded for treatment of pancreatic cancer</li> <li>– RHB-107 planned to be evaluated in combination with opaganib in an ongoing Phase 2a study in cholangiocarcinoma</li> </ul>
	<ul style="list-style-type: none"> <li>– <b>Ongoing U.S. Phase 2/3 study in non-hospitalized patients with symptomatic COVID-19 who do not require supplemental oxygen</b></li> </ul>

## RHB-107 - Ongoing Phase 2/3 COVID-19 Study

**Ongoing U.S. Phase 2/3 study with RHB-107 in non-hospitalized patients with symptomatic COVID-19 not requiring supplemental oxygen**

- ✓ **Study design:** 310 patients to be enrolled in a 2-part, multicenter, randomized, double-blind, placebo-controlled, parallel-group Phase 2/3 study
- ✓ **Innovative use of home-based monitoring technologies** with home nursing support allowing patients participate from home
- ✓ **Primary endpoint:** time to sustained recovery
- ✓ Patients will be tested for specific viral strain

**RHB-107 expected to be effective against emerging viral variants with mutations in the spike protein**

- ✓ Demonstrated potent inhibition of SARS-CoV-2 viral replication in in vitro model of human bronchial tissue
- ✓ Targets human cell factors involved in viral entry
- ✓ Simple once-daily orally-administered treatment





## **RHB-204**

*Investigational new drug*

*Targeting pulmonary nontuberculous Mycobacteria (NTM) disease - with QIDP designation, including Fast-Track development status*

***U.S. Phase 3 Study Ongoing***

# RHB-204 - Targeting First-Line Pulmonary NTM Disease

<b>Planned Indication</b>	<ul style="list-style-type: none"> <li>- Pulmonary Mycobacterium avium Complex (MAC) disease in adults with nodular bronchiectasis</li> </ul>
<b>The Product</b>	<ul style="list-style-type: none"> <li>- Patent-protected, oral all-in-one combination of three antibiotic drugs (clarithromycin, clofazimine and rifabutin) each known to be active against NTM disease caused by MAC<sup>i</sup></li> </ul>
<b>Key Attributes</b>	<ul style="list-style-type: none"> <li>- <b>Targeting first-line treatment - potential new standard-of-care for a disease with no FDA-approved first-line therapy</b></li> <li>- <b>Convenient stand-alone oral therapy for a chronic disease requiring extended treatment</b></li> <li>- Unique dosing combination - optimizing exposure for safety and efficacy</li> </ul>
<b>Market Size</b>	<ul style="list-style-type: none"> <li>- U.S. market potential estimated at approx. \$530M in 2021<sup>ii</sup></li> </ul>
<b>Development Status</b>	<ul style="list-style-type: none"> <li>- <b>Phase 3 study ongoing</b></li> <li>- QIDP and Fast-Track Designation granted, providing eligibility for rolling NDA review, Priority Review and accelerated approval</li> <li>- Orphan Drug designation extends potential U.S. market exclusivity to a total of 12 years post-approval</li> </ul>

<sup>i</sup> Wassilew et al, RESPIRATION 2021 <sup>ii</sup> Foster Rosenblatt

# RHB-204 for Pulmonary NTM - Background and Epidemiology

- **Difficult to treat infection with no FDA-approved first-line standard-of-care**
- NTM are a ubiquitous bacteria, mostly non-pathogenic but can cause human disease<sup>i</sup>
  - Pulmonary manifestations account for 80-90% of NTM associated disease<sup>ii</sup>
  - Approximately 80% of pulmonary NTM infections in the U.S. are associated with *Mycobacterium avium* complex (MAC)<sup>iii</sup>
- Pulmonary NTM disease symptoms can include fever, weight loss, chronic or recurring cough, chest pain, blood in sputum and fatigue<sup>iv</sup>
- NTM have high levels of drug resistance and require long term dosing with three or more antibiotics<sup>ii</sup>
- An orphan disease - with an estimated 110,000 pulmonary NTM patients in the U.S. in 2017<sup>v</sup>



<sup>i</sup> Wassilew et al, *RESPIRATION* 2016 <sup>ii</sup> Griffith DE, et al. An official ATS/IDSA statement: diagnosis, treatment, and prevention of nontuberculous mycobacterial diseases *Am J Respir Crit Care Med*. 2007;175(4):367-416; 3 American Thoracic Society; <sup>iii</sup> Prevots DR et al, *Am J Respir Crit Care Med* 2010; <sup>iv</sup> Daley et al *CHEST* 2017; <sup>v</sup> Foster Rosenblatt/Company estimates

## RHB-204 - CleaR-MAC Phase 3 Study Ongoing

**A Phase 3 study to assess RHB-204 as a first-line treatment for the Treatment of pulmonary Mycobacterium avium Complex (MAC) disease in adults with nodular bronchiectasis**

<b>Study Initiated</b>	<ul style="list-style-type: none"> <li>- November 2020</li> </ul>
<b>Study Design</b>	<ul style="list-style-type: none"> <li>- Multi-center, randomized, double-blind, two-part, placebo-controlled, parallel-group Phase 3 study; 3:2 randomization</li> <li>- Up to 40 U.S. clinical sites</li> <li>- Two-part study: <ul style="list-style-type: none"> <li>• Part one: placebo-controlled, subjects evaluated for primary endpoints at Month 6</li> <li>• Part two: Open-label treatment with RHB-204 for 10 months, with follow-up 3 months post-treatment</li> </ul> </li> </ul>
<b>Patient Population</b>	<ul style="list-style-type: none"> <li>- 125 subjects with symptomatic pulmonary MAC disease with nodular bronchiectasis</li> <li>- An interim sample size re-estimate is planned at approx. 50% enrolment</li> </ul>
<b>Endpoints</b>	<ul style="list-style-type: none"> <li>- Primary endpoints: <ul style="list-style-type: none"> <li>- Sputum culture conversion (SCC) at Month 6 of treatment</li> <li>- Quality of Life questionnaire – Bronchiectasis (QoL-B) respiratory symptoms domain score from Baseline to Month 6</li> </ul> </li> </ul>



## RHB-104

*Investigational new drug*

*Combination therapy targeting MAP bacteria for treatment of Crohn's disease and potentially other autoimmune diseases*

***Positive First Phase 3 Study in Crohn's Disease -  
Study Successfully Met Its Primary Endpoint and Key Secondary Endpoints***

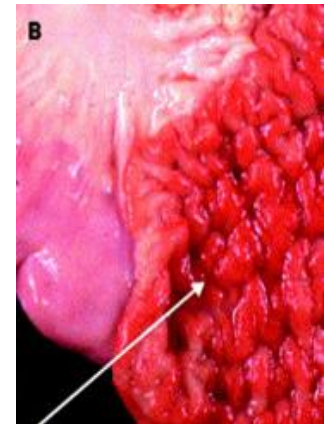
# The Link between Crohn's Disease and MAP

## Growing evidence that intracellular mycobacteria play a crucial role in Crohn's disease

- MAP (*Mycobacterium avium* subsp. *paratuberculosis*) is the causative agent of Johne's disease, an infectious disease in cattle, clinically and pathologically similar to Crohn's disease
  - An intracellular pathogen that proliferates in monocytes/macrophages
  - Extremely slow growing and widely pervasive in the environment
- Advances in diagnostic technology have led to increasingly higher identification of MAP in Crohn's diseases patients
  - 92% (34/37 Crohn's disease patients by PCR) - Bull, J Clin Microbiol, 2003
  - 86% (52/60 Crohn's disease patients) - Shafran, Dig Dis Sci, 2002
- Crohn's disease is a multifactorial disease
  - Defective innate immunity to intracellular bacteria
  - Mutations in the NOD2 gene are strongly associated with Crohn's disease
- Mycobacterial infections in humans are difficult to treat; Effective anti-mycobacterial agents require intracellular activity
  - ATS/IDSA<sup>i</sup> and WHO advise triple antibiotic therapy for non-tuberculosis mycobacterial disease



Crohn's disease



Johne's disease

<sup>i</sup> American Thoracic Society, Infectious Disease Society of America, World Health Organization

## RHB-104 for Crohn's - Product and Market Overview

<b>Planned Indication</b>	Treatment of Crohn's disease in adult patients
<b>The product</b>	Patent-protected combination of 3 antibiotics (clarithromycin, clofazimine and rifabutin) in a single oral capsule with potent intracellular, antimycobacterial and anti-inflammatory properties
<b>Market Size</b>	Worldwide market estimated to exceed \$12.8 billion in 2022 <sup>i</sup>
<b>Positive First Phase 3 Top-Line Results</b>	Positive results from the MAP US first Phase 3 study with RHB-104 in Crohn's disease - study successfully met its primary endpoint and key secondary endpoints
<b>Key Attributes</b>	<ul style="list-style-type: none"> <li>✓ Unique MoAs, including targeting the potential underlying cause of Crohn's</li> <li>✓ Improved efficacy on top of standard-of-care</li> <li>✓ Safety and tolerability enabling combining RHB-104 with standard-of-care</li> <li>✓ Oral all-in-one capsule</li> </ul>

<sup>i</sup> GlobalData Crohn's Disease Global Forecast, September 2017

# RHB-104 Positive First Phase 3 Study in Crohn's - Met Primary Endpoint and Key Secondary Endpoints

Multi-center, randomized, double-blind, placebo-controlled, parallel group study (MAP US study) to assess efficacy and safety of orally-administered, fixed-dose combination RHB-104 in subjects with moderately to severely active Crohn's disease, on-top of baseline background SoC medications

<b>Number of Subjects</b>	331
<b>Sites</b>	Over 100 in the U.S., Canada, Europe, Australia, New Zealand and Israel
<b>Primary Endpoint</b>	State of remission at week 26
<b>Previous Studies</b>	<ul style="list-style-type: none"> <li>- Several PK, Phase 1, 2 and 3 clinical studies conducted</li> </ul>
<b>Phase 3 MAP US Development Status</b>	<ul style="list-style-type: none"> <li>- Positive Phase 3 results - study met its primary endpoint and key secondary endpoints</li> <li>- Ongoing discussions with KOLs FDA meeting planned to discuss design of confirmatory Phase 3 study and path to potential approval</li> </ul>



# RHB-104 Crohn's Disease Phase 3 - Positive Results

The MAP US Phase 3 study met its primary and key secondary endpoints, demonstrating meaningful, consistent and statistically significant clinical activity of orally-administered RHB-104 as an add-on to standard-of-care treatments

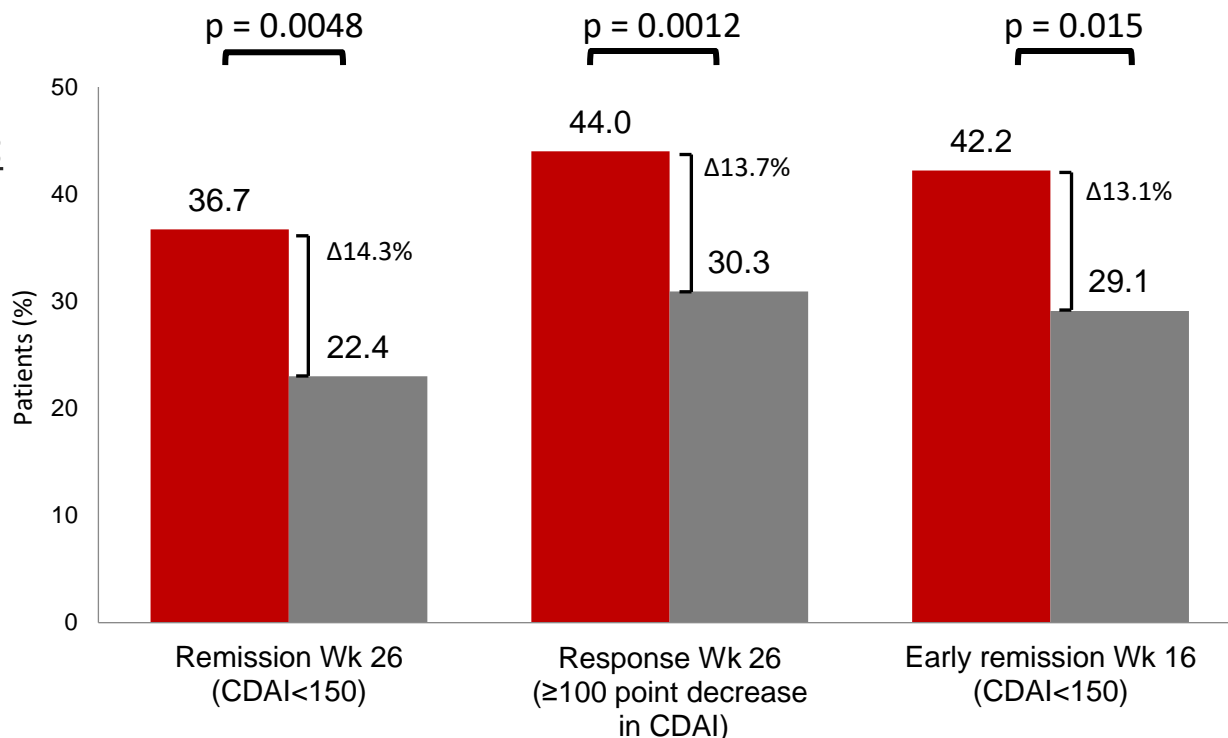
■ RHB-104 (n= 166)    ■ Placebo (n=165)

## Primary Endpoint Met:

- ✓ Remission at week 26

## Secondary Endpoints Met:

- ✓ Response at week 26
- ✓ Early remission at week 16



<sup>i</sup> Calculated with Cochran-Mantel-Haenszel (CMH) chi-square test with stratification according to anti-TNF agents use (yes/no)

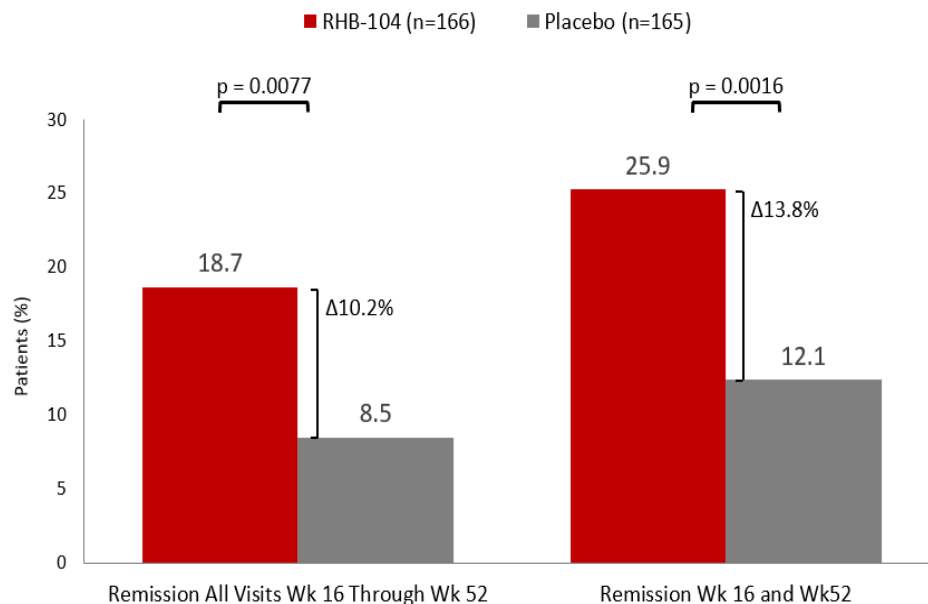
# RHB-104 Crohn's Disease Phase 3 - Positive Results

## Additional Endpoints Met:

- ✓ **Durable remission weeks 16-52:** statistically significant benefit in durable remission on all visits, weeks 16-52, demonstrating an improvement of 100% over placebo
- ✓ **Maintenance of remission:** analysis of maintenance of remission at week 52 in subjects noted to be in remission at week 16 demonstrated statistically significant benefit with RHB-104 over placebo

## Safety:

- ✓ **RHB-104 was demonstrated to be generally safe and well-tolerated** similar TAEs, SAEs and AEs leading to study drug discontinuation between treatment groups
- ✓ ECG monitoring report demonstrated evidence of progressive prolongation of the QTcF interval across visits; None of these QT abnormalities resulted in adverse cardiac events

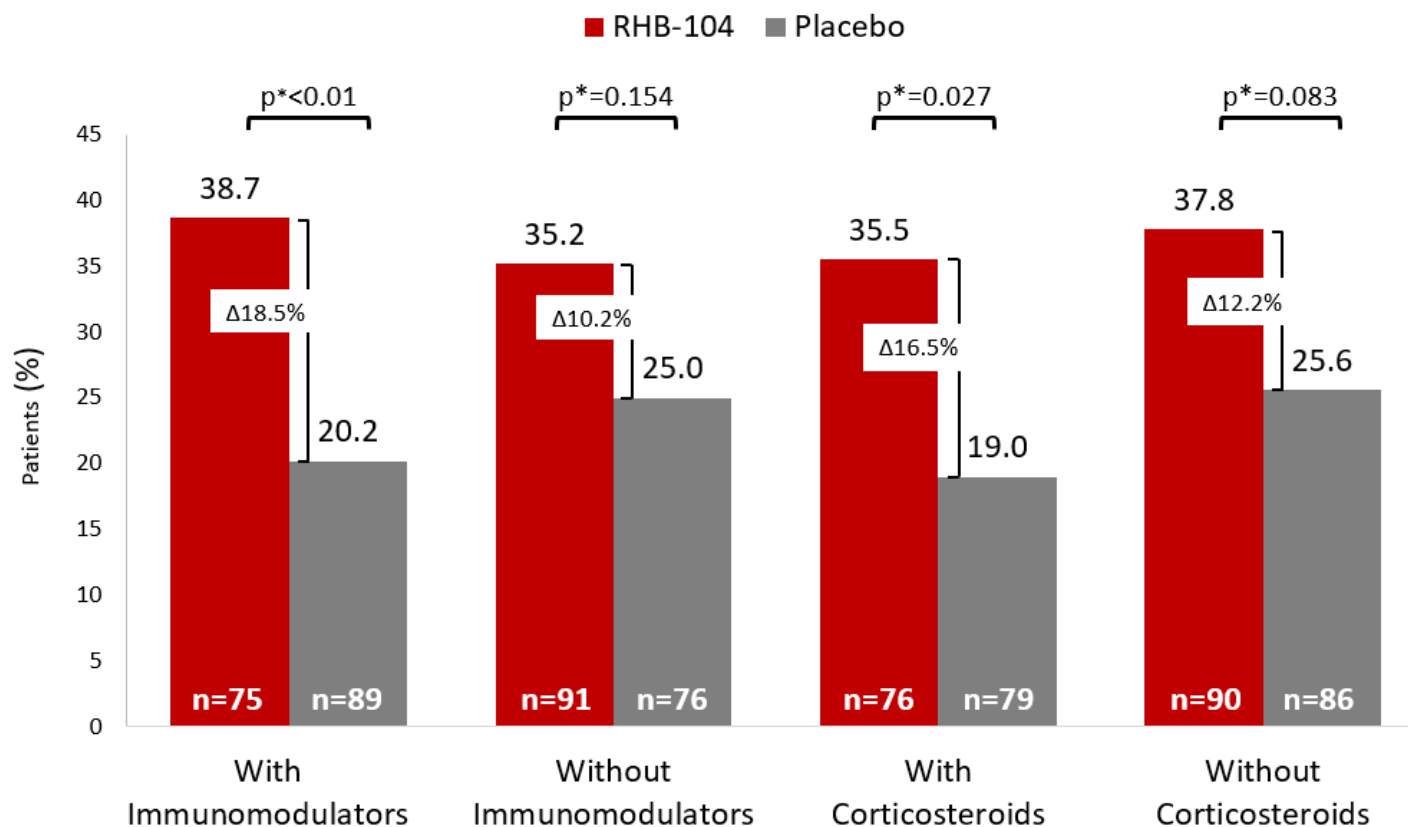


<sup>i</sup> Calculated with Cochran-Mantel-Haenszel (CMH) chi-square test with stratification according to anti-TNF agents use (yes/no); <sup>o</sup> Number of subjects reflects those subjects who have completed week 52 assessments and are no longer receiving blinded therapy

## RHB-104 Crohn's Disease Phase 3 - Positive Results

**Effects of concomitant immunomodulators and corticosteroids:** Remission at week 26 concomitant meds subgroup analyses demonstrated the impact of RHB-104 as an add-on therapy to a variety of standard-of-care agents despite not being prospectively powered

- ✓ Meaningful and statistically significant treatment effects were observed in patients using concomitant immunomodulators (38.7% vs. 20.2%,  $p < 0.01$ ) and corticosteroids (35.5% vs. 19.0%,  $p = 0.027$ ) throughout the trial

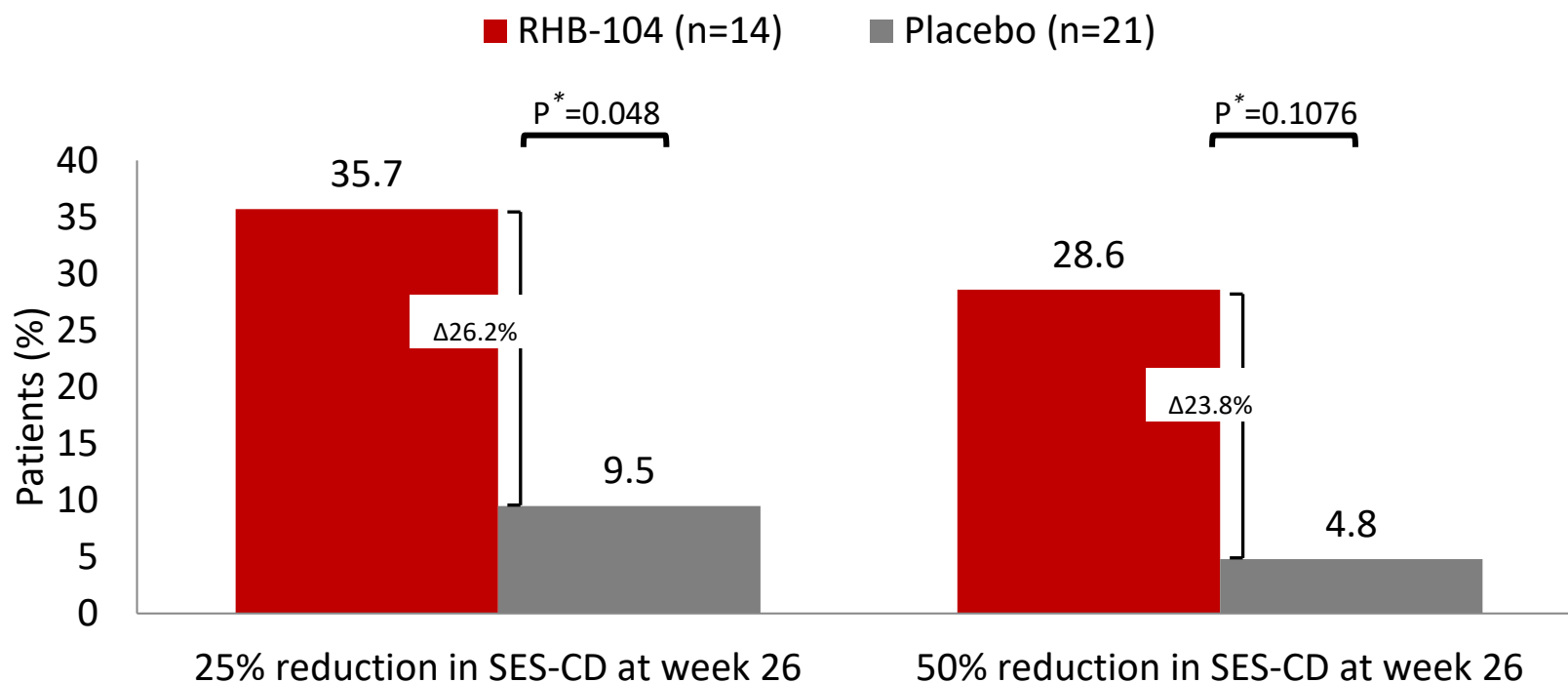


<sup>i</sup> Calculated with Cochran-Mantel-Haenszel (MH) chi-square test with stratification according to anti-TNF agent use (yes/no); <sup>o</sup> Calculated with Mantel-Haenszel (MH) chi-square test

# RHB-104 Crohn's Disease Phase 3 - Positive Results

## Endoscopic response:

- ✓ Endoscopy was voluntary in the study
- ✓ In a small subset of patients (n=35) in whom endoscopy was performed, results showed meaningful improvement in endoscopic healing (Simple Endoscopic Score for Crohn's Disease (SES-CD)) at week 26 (statistically significant at 25% decrease with similar trend at 50% decrease)

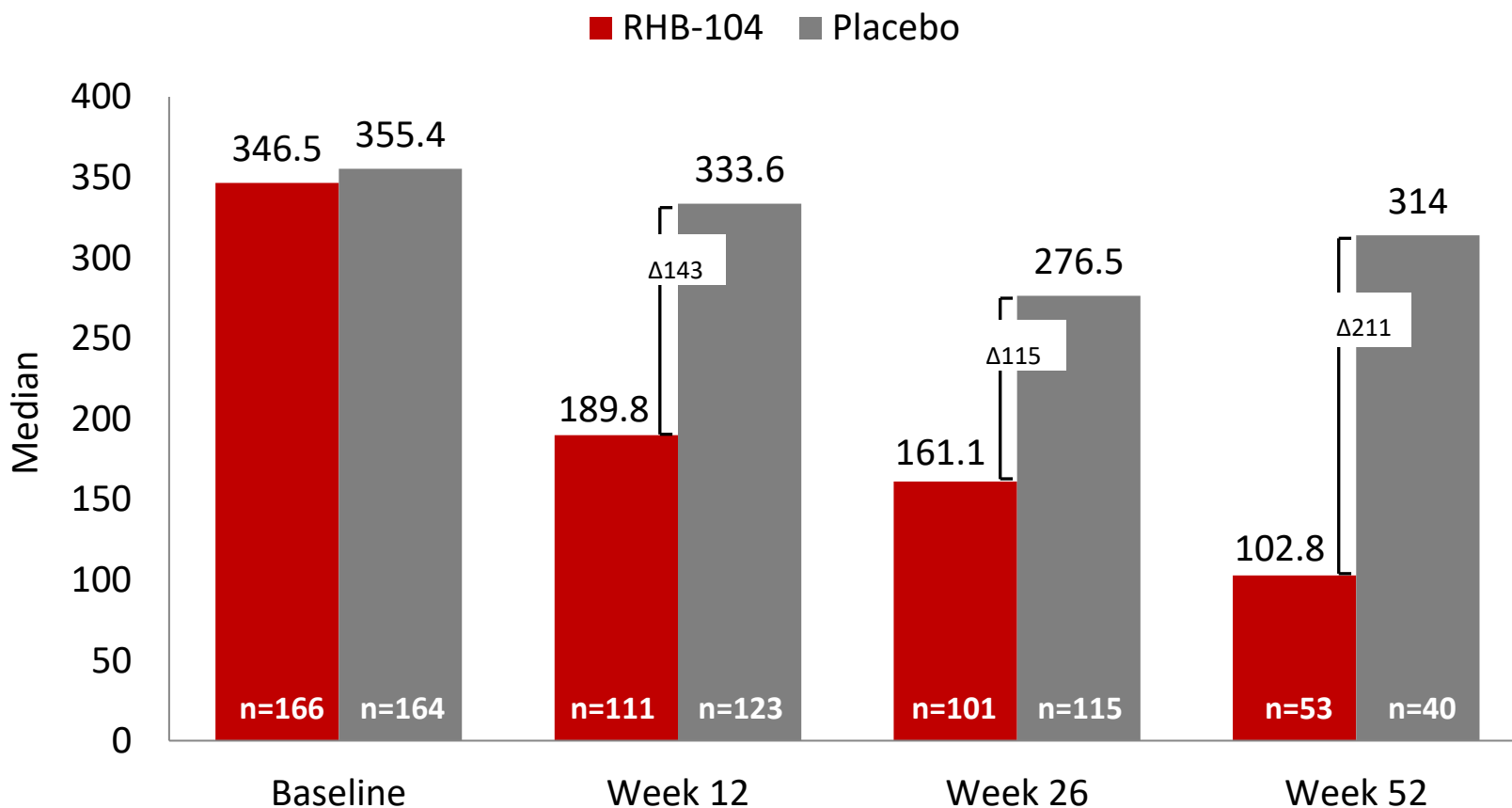


<sup>i</sup> Calculated with Cochran-Mantel-Haenszel (CMH) chi-square test with stratification according to anti-TNF agents use (yes/no)

# RHB-104 Crohn's Disease Phase 3 - Positive Results

## Median Fecal Calprotectin

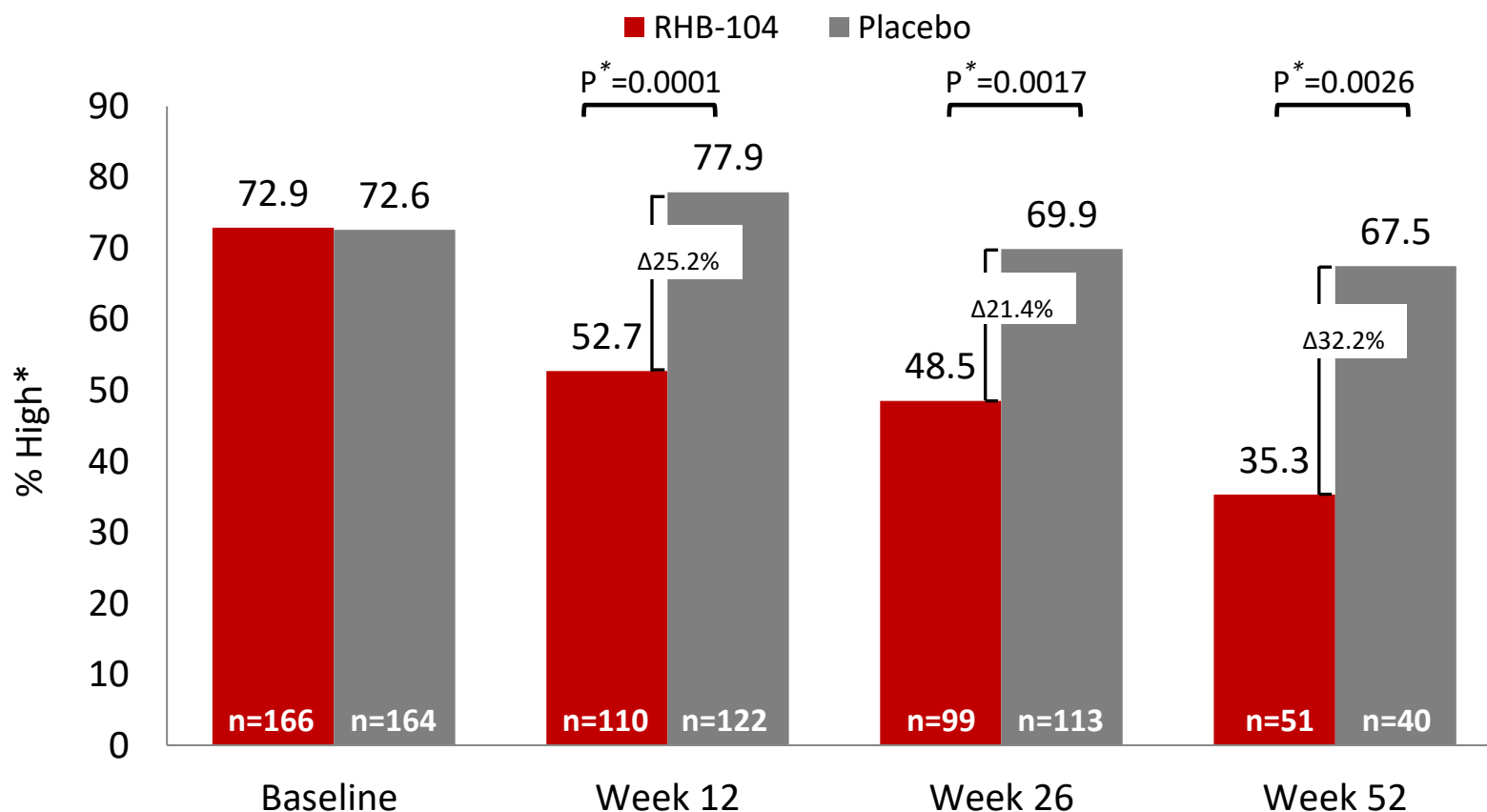
- ✓ Marked and continuous decline of calprotectin inflammatory marker in the RHB-104 active arm from baseline through week 52



# RHB-104 Crohn's Disease Phase 3 - Positive Results

## % High Fecal Calprotectin\*\*

- ✓ Proportion of high calprotectin declined significantly in the RHB-104 active arm from baseline through week 52 (weeks 12, 26 and 52)



<sup>i</sup> Calculated with Cochran-Mantel-Haenszel (CMH) chi-square test with stratification according to anti-TNF agents use (yes/no); \*\* High defined as > 162.9

## RHB-104 - Multiple Barriers to Entry

- Robust global patent portfolio covering RHB-104, with additional claims being pursued
- Potential 3 years exclusivity for treatment of Crohn's disease
- Limited availability of clofazimine (distributed by the World Health Organization (WHO) and Novartis) and generally requires name based individual import permit or Expanded Access Program (EAP) process for use
- Lower concentrations of the triple combination of RHB-104 active components provide excellent synergistic anti-MAP growth activity compared to individual or dual combinations of the drugs<sup>i</sup>
- All-in-one capsule solution for patients and physicians (reduced co-pays, etc.)
- Potential PK synergies of APIs administered in the RHB-104 formulation that disappear when administered concomitantly
- APIs are not available in doses being used in RHB-104
- Physicians' potential liability exposure and complicated ramp-up period

<sup>i</sup> Alcedo KP, Thanigachalam S, Naser SA. RHB-104 triple antibiotics combination in culture is bactericidal and should be effective for treatment of Crohn's disease associated with *Mycobacterium paratuberculosis*, *Gut Pathogens*, 2016, 8:32.



## **RHB-102<sup>i</sup>** *Investigational new drug*

*A bi-modal extended release, once-daily, ondansetron*

***RHB-102 24 mg - Positive Results from a U.S. Phase 3 Study for Gastroenteritis/Gastritis - Met Primary Endpoint***

***RHB-102 12 mg - Positive Results from a U.S. Phase 2 Study for IBS-D Met Primary Endpoint***

<sup>i</sup> Bekinda® is the proposed tradename for RHB-102, which is subject to review by the FDA at the time of NDA filing



# RHB-102 24 mg: Gastroenteritis & Gastritis - Positive First Phase 3 Results

A Randomized, Double-Blind, Placebo Controlled, Parallel Group Phase 3 Study (GUARD study) to Assess the Safety and Efficacy of RHB-102<sup>i</sup> 24 mg for the Treatment of Acute Gastroenteritis & Gastritis

<b>The Product</b>	Patent-protected, once-daily extended release oral tablet ondansetron 24 mg
<b>Number of Subjects</b>	321 adults and children over the age of 12
<b>Sites</b>	21 in the U.S.
<b>Primary Endpoint</b>	The absence of vomiting, without rescue medication and intravenous hydration, from 30 minutes post first dose of the study drug until 24 hours post dose
<b>Potential Advantages</b>	<ul style="list-style-type: none"> <li>- If approved by FDA, RHB-102 could become the first 5-HT<sub>3</sub> antiemetic drug indicated for the treatment of acute gastroenteritis or gastritis in the U.S.</li> <li>- Long-lasting (24H) oral treatment with the potential to reduce dehydration and hospital visits and stays</li> </ul>
<b>Market Size</b>	<ul style="list-style-type: none"> <li>- Approximately 179 million cases of acute gastroenteritis annually in the U.S., leading to an estimated 470,000 hospitalizations<sup>ii</sup></li> <li>- Worldwide potential market could exceed \$650 million annually<sup>iii</sup></li> </ul>
<b>Development Status</b>	<ul style="list-style-type: none"> <li>- <b>Positive Phase 3 study results announced June 2017 - the study met primary endpoint</b></li> <li>- <b>RedHill designing a confirmatory Phase 3 study for acute gastroenteritis and gastritis</b></li> </ul>

<sup>i</sup> Bekinda® is a proposed tradename for RHB-102, which is subject to FDA review and approval

<sup>ii</sup> Scallan E, Griffin PM, Angulo FJ, Tauxe RV, Hoekstra RM. Foodborne Illness Acquired in the United States - Unspecified Agents. *Emerg Infect Dis.* 2011;17(1):16-22

<sup>iii</sup> Graves S. Nancy, Acute Gastroenteritis, *Prim Care Clin Office Pract* 40 (2013) 727–741 and Company analysis;

# RHB-102 24 mg: Gastroenteritis & Gastritis Positive Phase 3 Results

A Randomized, Double-Blind, Placebo Controlled, Parallel Group Phase 3 Study (“GUARD”) to Assess Safety and Efficacy of RHB-102 24 mg for Treatment of Acute Gastroenteritis & Gastritis

## Results

- **The Phase 3 GUARD study successfully met its primary endpoint in the Intent to Treat (ITT) population ( $p = 0.04$ )**
- **ITT:** RHB-102 24 mg improved the efficacy outcome by 21%; 65.6% of RHB-102 24 mg treated patients as compared to 54.3% of placebo patients ( $p = 0.04$ ;  $n=192$  in the RHB-102 group and  $n=129$  in the placebo group)
- **PP:** In patients who met all protocol entry criteria and for which the diagnosis of gastroenteritis was confirmed ( $n=177$  in the RHB-102 group and  $n=122$  in the placebo group), RHB-102 24 mg improved the efficacy outcome by 27%; 69.5% of patients in the RHB-102 24 mg group vs. 54.9% in the placebo group, ( $p = 0.01$ )
- RHB-102 24 mg was demonstrated to be safe and well-tolerated; electrocardiogram results showed no adverse changes with treatment

# RHB-102 12 mg: IBS-D

## Positive Phase 2 Results

A Randomized, Double-Blind, Placebo-Controlled, 2-Arm Parallel Group Phase 2 Clinical Study Designed to Evaluate the Safety and Efficacy of RHB-102 12 mg in Patients Suffering from Diarrhea-Predominant Irritable Bowel Syndrome (IBS-D)

<b>The Product</b>	Patent-protected, once-daily extended release oral tablet ondansetron 12 mg
<b>Number of Subjects</b>	126 patients
<b>Sites</b>	16 in the U.S.
<b>Primary Endpoint</b>	Response in stool consistency compared to baseline, per FDA guidance definition
<b>Market Size</b>	<ul style="list-style-type: none"> <li>- It is estimated that at least 30 million Americans may suffer from IBS, of which approximately 40% are of the IBS-D subtype<sup>i</sup></li> <li>- 8.3 million diagnosed prevalent cases in 2018 in the U.S.<sup>ii</sup></li> <li>- 2019 potential U.S. market for IBS-D treatments estimated to reach ~ \$980M</li> </ul>
<b>Development Status</b>	<ul style="list-style-type: none"> <li>- <b>Positive top-line results announced Oct. 2017; Study successfully met primary endpoint</b></li> <li>- <b>Positive End-of-Phase 2/Pre-Phase 3 FDA meeting announced Sep. 2018 - design of two pivotal Phase 3 studies with RHB-102 to be finalized and partnership discussions accelerated</b></li> </ul>

<sup>i</sup> Lovell RM, Global prevalence of and risk factors for irritable bowel syndrome: a meta-analysis, *Clin Gastroenterol Hepatol* (2012), 10(7)712-721; Saito YA et al, The epidemiology of irritable bowel syndrome in North America: a systemic review, *Am J Gastroenterol* (2002), 97(8): 1910-5; <sup>ii</sup> GlobalData

# RHB-102 12 mg: IBS-D

## Positive Phase 2 Results

A Randomized, Double-Blind, Placebo-Controlled, 2-Arm Parallel Group Phase 2 Clinical Study Designed to Evaluate the Safety and Efficacy of RHB-102 12 mg in Patients Suffering from Diarrhea-Predominant Irritable Bowel Syndrome (IBS-D)

### Results

- **Primary endpoint:** The Phase 2 study met its primary endpoint, improving the primary efficacy outcome of stool consistency response (per FDA guidance definition) by an absolute difference of 20.7% ( $p=0.036$ )
  - 56.0% responders of subjects treated with RHB-102 ( $n=75$ ) vs. 35.3% responders of the placebo subjects ( $n=51$ )
- **Secondary endpoints:** While not powered for statistical significance of the secondary efficacy endpoints, the study suggested clinically meaningful improvement in both abdominal pain response and overall response (combined stool consistency and abdominal pain response)
- **Safety:** RHB-102 12 mg was demonstrated to be safe and well-tolerated



## **RHB-106**

*Investigational new drug*

*Proprietary Encapsulated Formulation for Bowel Preparation*

## RHB-106 - Encapsulated Bowel Preparation

<b>Planned Indication</b>	Preparation of the gastrointestinal tract prior to abdominal procedures/surgeries, such as colonoscopy
<b>The Product</b>	<ul style="list-style-type: none"> <li>– Proprietary, flavorless, odorless, solid oral encapsulated formulation for bowel preparation</li> <li>– Caplet formulation composed of sodium picosulfate, magnesium oxide, ascorbic acid and simethicone, mannitol and other excipients</li> </ul>
<b>Potential Advantages</b>	<ul style="list-style-type: none"> <li>– Improved safety over existing encapsulated preparations</li> <li>– Convenient, easy, taste free, and odorless</li> <li>– Can be taken with water</li> </ul>
<b>Potential Market Size</b>	<ul style="list-style-type: none"> <li>– Approx. 19 million colonoscopies performed annually in the U.S<sup>i</sup></li> <li>– 2019 U.S. sales of bowel preparations estimated at approximately \$580 million<sup>ii</sup></li> </ul>
<b>Development Status</b>	<ul style="list-style-type: none"> <li>– Phase 2/3 studies planned</li> <li>– Phase 2a with a previous formulation of RHB-106 conducted in 62 patients in Australia<sup>iii</sup> demonstrating significantly improved patient response and comparable bowel cleansing</li> </ul>

<sup>i</sup> IDATA Research, August 2018; <sup>ii</sup> Foster-Rosenblatt, December 2019; <sup>iii</sup> Ramrakha SR et al. Investigating the tolerability of four bowel preparations – the taste test, J Gastroenterol Hepatol, 2006; 21 (Suppl;4): A286-A299



**Thank You!**

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