

Redeye Investor Forum Online

February 24th, 2022 Anna Ljung, CEO



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Potential new global market leader in Onychomycosis

MOB-015 has demonstrated world-leading ability to kill nail fungus

 76%¹ of patients became fungus free, in two phase 3-studies including 800+ patients

- Additional de-risked US phase 3 study based on completed phase 3 studies to enable US approval and strengthen claims globally
- Targeting category leadership with USD 250-500m potential global product sales







Republic of Korea

Japan

FU



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Partnerships in place – potential

milestones of USD 120m

Scandinavia

Canada

 Swedish MPA reference country for EU submission March 2022
Product launch expected 2023

On track for launch – capturing full

value potential

- Proven commercial track record from Kerasal Nail[®] – built SEK 440 million franchise with 30% market share in the US
- Commercialization process to be repeated for MOB-015



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100+ million patients need better treatment in EU/US only

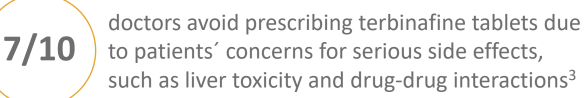


of the population suffer from nail fungus¹



10%

global onychomycosis market²



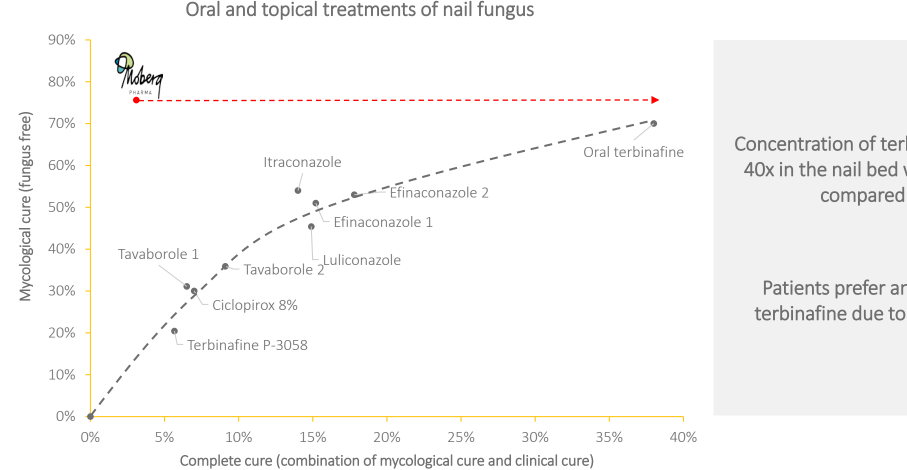


1) PLoS Pathog. 2014 Jun; 10(6): e1004105.

2) Moberg Pharma estimate, based on market data from Symphony Health Solutions (US Rx sales), Symphony IRI (US OTC sales), and market data from Moberg Pharma's partners.

3) LifeSci Physician Survey, April 4, 2017

Superior mycological cure – expecting to increase complete cure



Concentration of terbinafine 1000x in the nail and 40x in the nail bed when treated with MOB-015 compared to oral terbinafine.

Patients prefer an efficacious topical to oral terbinafine due to risk for severe side effects.

Additional Phase 3 study with attractive commercial impact



Shorter dosing regimen

Patient benefit

Competitive advantage

- A regimen with daily dosing for 8-12 weeks followed by once weekly treatment, is highly attractive, and expected to maintain high mycological cure to deliver high complete cure
- This will significantly strengthen claims globally for MOB-015



- Shorter daily dosing for only 8-12 weeks would be a significant improvement for patients, leading to improved convenience and compliance
- 75% of patients see improvements already at week 12¹

- Main topical competitors have 48 weeks daily treatment, but poor compliance. Average consumption is 12-16 weeks²
- MOB-015's dosing regiment will compare to oral treatment but without the safety issues of oral treatments



Key Opinion Leaders strongly support the concept



Dr Boni Elewski Professor and Chair of the Department of Dermatology University of Alabama "The high mycological cure rate demonstrated is very impressive and given the rapid onset of the antifungal effect, MOB-015 offers exciting benefits. I will definitely use it for my patients. A higher complete cure rate is likely to be achieved with a shorter treatment period and this would also be much more attractive to patients."







Dr Aditya Gupta Professor, Department of Medicine University of Toronto *"I am a strong supporter of this concept*. With an optimized dosing regimen this product has *great potential* and *may become the preferred therapeutic option*, not only for monotherapy, but also as maintenance therapy to reduce recurrence after oral treatment."





Dr Jan Faergemann Professor in Dermatology Sahlgrenska Academy University of Gothenburg "Based on decades of experience with terbinafine and the excipients used in MOB-015, I believe a shorter treatment period has the potential to provide higher complete cure rates. *Killing the fungus is the driver of also reaching complete cure.*"



Preparations ongoing for the next Ph3-study in U.S.

- Similar design as the already completed North American study
- Multi-center, double-blind, randomized, vehicle-controlled study
- Scheduled to include 350 patients in North America
- Purpose of the new study:

- Enable market approval in the U.S.
- Strengthen the product's clinical data and marketing claims globally
- The risk in the new study is significantly reduced through the experience gained from the previous studies
 - Cooperation with the same CRO and lead investigator as in the previous North American study
- Goal to submit documentation on the new study to the FDA and ethical committee in Q1 2022

Swedish MPA will be reference member state – expected approval and launch 2023



- The Medical Products Agency in Sweden has agreed to be reference member state for Moberg Pharma AB's registration application for MOB-015
 - the Swedish Medical Products Agency has announced that the application can be submitted in March 2022.
 - Moberg Pharma will submit a full application, which offers the possibility of data exclusivity in Europe for up to 10 years following market approval.
 - The company will submit the registration application in Europe through the decentralized process, and market approval is expected in 2023.
- EMA's Paediatric Committee approval (September 2021) paves way for EU submission
 - Supplementary pediatric study during and after approval process for MOB-015



US USD 150 - 300m	US Rx¹ potential : USD 150 - 300m (400 - 600 thousand units à USD 375 - 500/unit after GTN discount i.e. pricing on par with branded competitors and a target market share of 8 - 12%)
Other Rx markets	Other Rx markets, e.g. Japan and Canada:
USD 50 - 100m	USD 50 - 100m (USD 40 - 100/unit ex factory and targeting a market share of 10 - 20%)
OTC markets	OTC markets in EU and RoW :
USD 50 - 100m	USD 50 - 100m (3.5 - 7 million units à EUR 15/unit ex factory)

Strong commercial partners in place



- The world leader in OTC antifungal treatments with the brand Canesten
 - Up to EUR 50 million in milestone payments with EUR 1.5 million upfront
 - Royalties and supply fees for delivered products

USD 290m Japanese market for branded

USD 200m+

EU OTC market for topical

onychomycosis in 2017

drugs for onychomycosis in 2018



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- Up to USD 50 million in milestone payments, with USD 5 million upfront
- Royalties and supply fees for delivered products

USD 58m

Canadian market for onychomycosis prescription drugs in 2017

- Up to USD 14.6 million in milestone payments, with USD 0.5 million upfront
- Royalties on future net sales in Canada

USD 40m Korean market for topical drugs

n market for topical drugs for onychomycosis



- Korean dermatology market leader, excellent coverage of dermatology clinics
- Distribution agreements with attractive margins



Scandinavian OTC market for topical onychomycosis

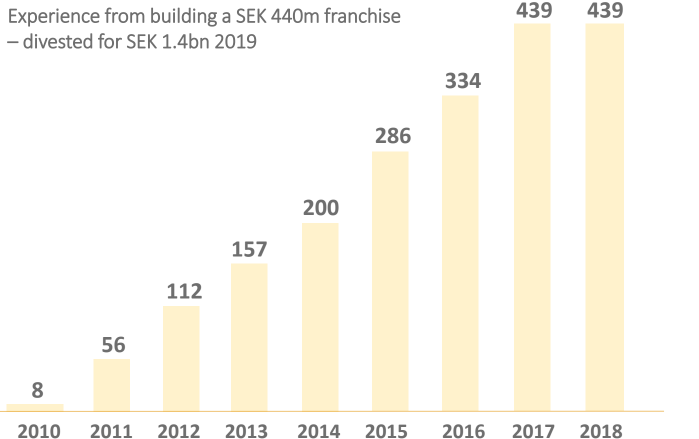


 Allderma is managed by the team responsible for the successful Nordic launch of Nalox[®], our first-generation nail fungus product

Excellent commercial track record from Kerasal Nail

Net Sales, SEKm

Experience from building a SEK 440m franchise



Proven commercial track record with leading OTC brand Kerasal Nail[®] for nail fungus

- Distributors in **30+ markets**
- Direct sales in the U.S. with #1 position, 30% market share and available in more than 30,000 U.S. stores

Commercialization process to be repeated for MOB-015

- Focus on podiatrists: >40% US prescriptions
- DTC marketing to U.S. consumers ۲
- Co-promotion with U.S. derm company

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