



Medicines for Rare Skin Diseases and Conditions –
A Gene Therapy Company

C O R P O R A T E P R E S E N T A T I O N
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SFH-0187556

Forward-Looking Statements

This presentation contains forward-looking statements and information. The use of words such as “may,” “might,” “will,” “should,” “expect,” “plan,” “anticipate,” “believe,” “estimate,” “project,” “intend,” “future,” “potential,” or “continue,” and other similar expressions are intended to identify forward-looking statements. These forward-looking statements are predictions and involve risks and uncertainties such that actual results may differ materially from these statements. These risks and uncertainties include, among others: actions by the FDA and other regulatory agencies, results and timing of current and planned clinical trials, risks related to the commercialization of our products, our ability to manufacture sufficient quantities of products for clinical trials and commercial launch, and those other risks detailed from time to time under the caption “Risk Factors” and elsewhere in Krystal Biotech's Securities and Exchange Commission (SEC) filings included in our Annual Report on Form 10-K for the year ending December 31, 2017, and in future filings and reports of Krystal Biotech. The Company undertakes no duty or obligation to update any forward-looking statements as a result of new information, future events or changes in its expectations.

Company Overview

- NASDAQ: KRYS; Started operations in 2016 with headquarters in Pittsburgh, Pennsylvania.
- Established a proprietary fully-integrated HSV-1-based gene therapy platform and a pipeline of clinical and non-clinical effectors to target rare diseases and conditions. Zero royalty burden.
- Interim data readout in GEM-Phase I/II trial targeting Dystrophic Epidermolysis Bullosa (DEB) met all primary and secondary endpoints.
 - Data readout anticipated in 1H 2019
 - Pivotal study anticipated to begin in 2H 2019
 - BLA filing expected in 1H 2020
- Multiple targets (5 additional indications) based on platform currently in pipeline.
- US Patents 9,877,990 (issued 1/16/18) and 10,155,016 (issued 12/18/18) covering pharmaceutical compositions and methods of their use.
- First GMP In-house manufacturing facility in Pittsburgh, PA complete. Plans to build a second GMP facility in 2H 2019 in motion.
- Insider ownership (management, employees, directors): 34 percent of fully diluted shares outstanding (as of 3/31/19)

Fully-Integrated Vector Platform

Modified Herpes Simplex Virus 1 (HSV-1) vector well suited to treat skin diseases

Proprietary Vectors

and underlying cell lines support robust and flexible drug production

Direct delivery

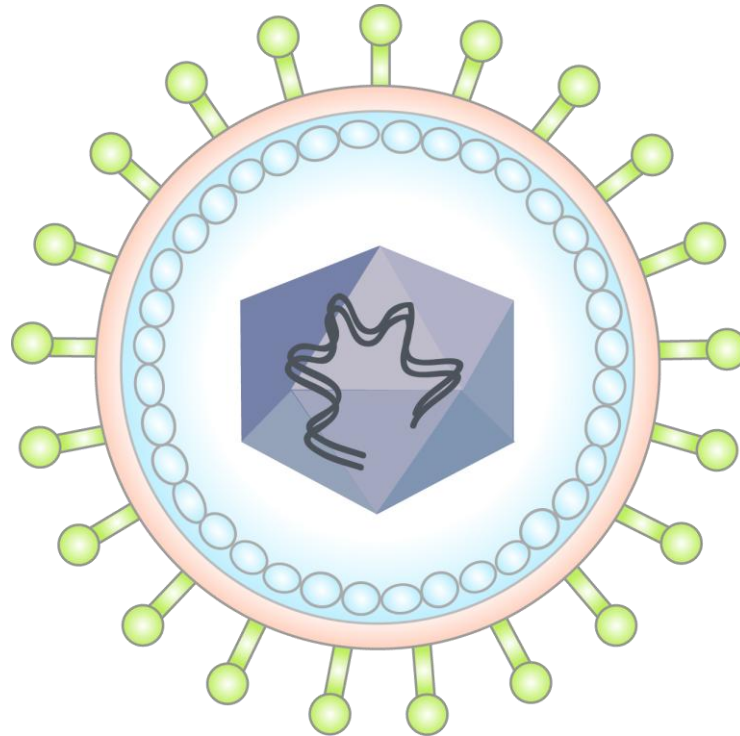
Topical administration for open wounds and intradermal for intact skin

Reproducible and Scalable Manufacturing

using internally developed and validated protocols

Non-integrating

into the DNA making it safer



Significant payload capacity

due to ~150Kb genome to accommodate multiple genes and effectors in the backbone

Stability

of vector beneficial to production and storage

High Transduction Efficiency

Transduces dividing and non-dividing skin cells

Non-Replicating

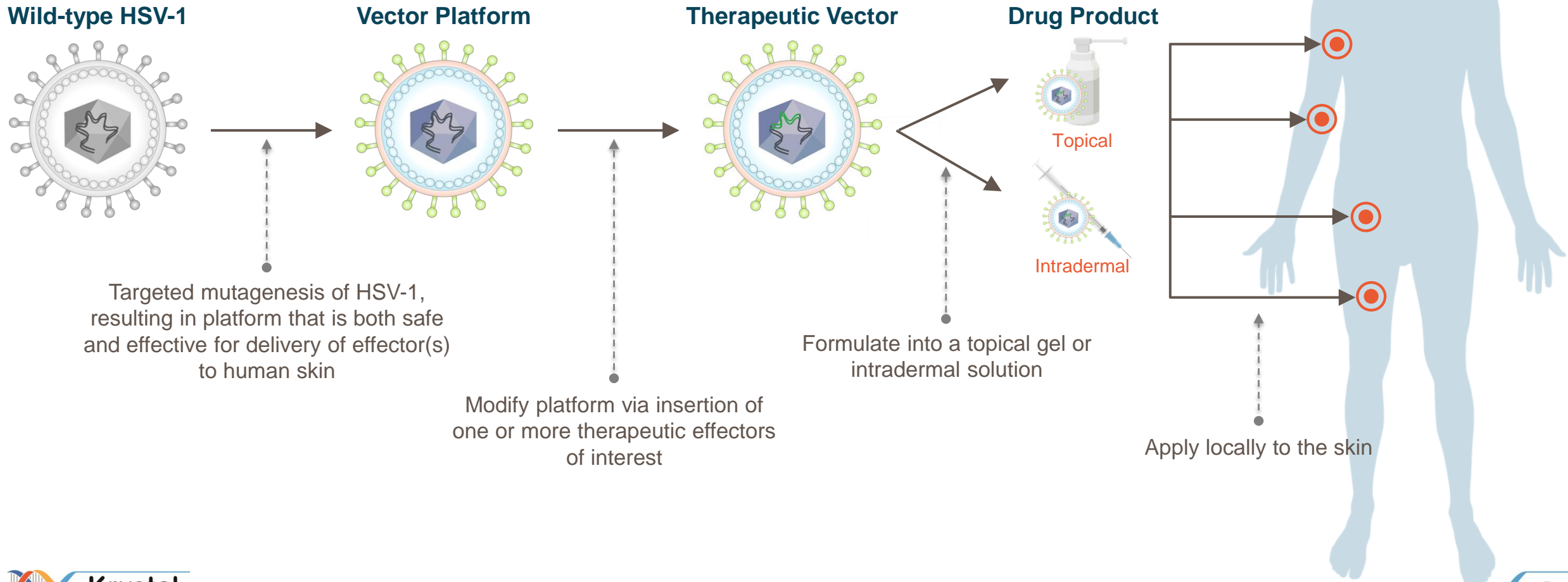
Safe for repeat administration; transient transgene expression, diluted by cell divisions

Regulatory precedent

HSV-1 used as backbone in Amgen's Imlytic[®], which is approved for melanoma and administered weekly to patients

Krystal's Unique and Straightforward Approach

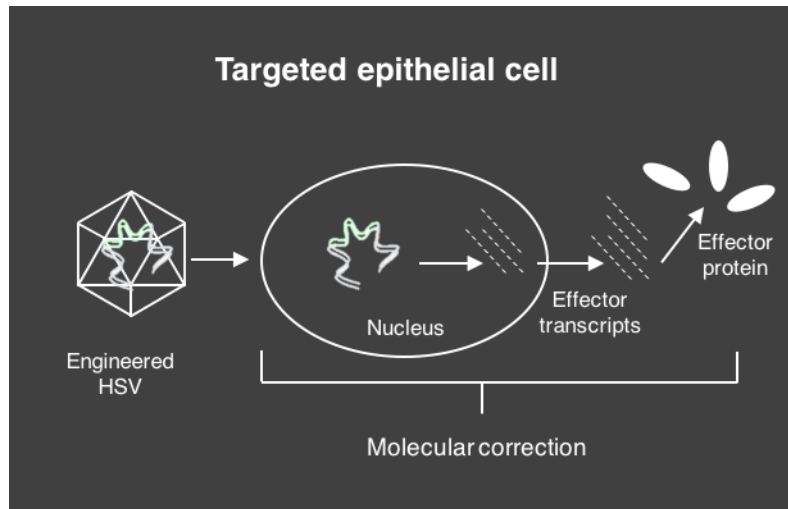
“Off-the-shelf” gene therapy with repeat administration



Krystal's Approach: Applicable to a Wide Range of Skin Diseases and Conditions

1. Severe monogenic skin diseases

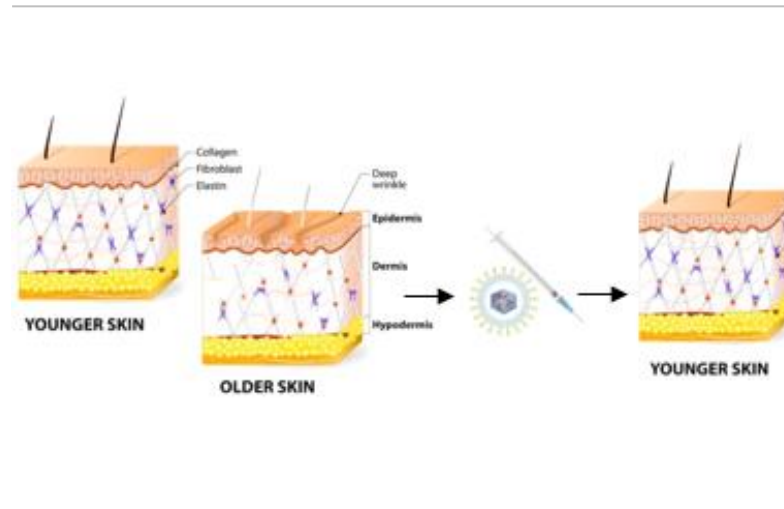
Indications: Dystrophic Epidermolysis Bullosa (DEB); Autosomal Recessive Congenital Ichthyosis (ARCI); Junctional Epidermolysis Bullosa (JEB); Netherton Syndrome (NS)



Effectors: *COL7A1* (type VII collagen); *TGM1* (transglutaminase-1); laminins; *SPINK5* (serine protease inhibitor kazal-type 5)

2. Aesthetic conditions

Indications: fine lines; nasolabial folds; glabellar lines; depressed scars; UV-induced skin damage



Effectors: Collagens

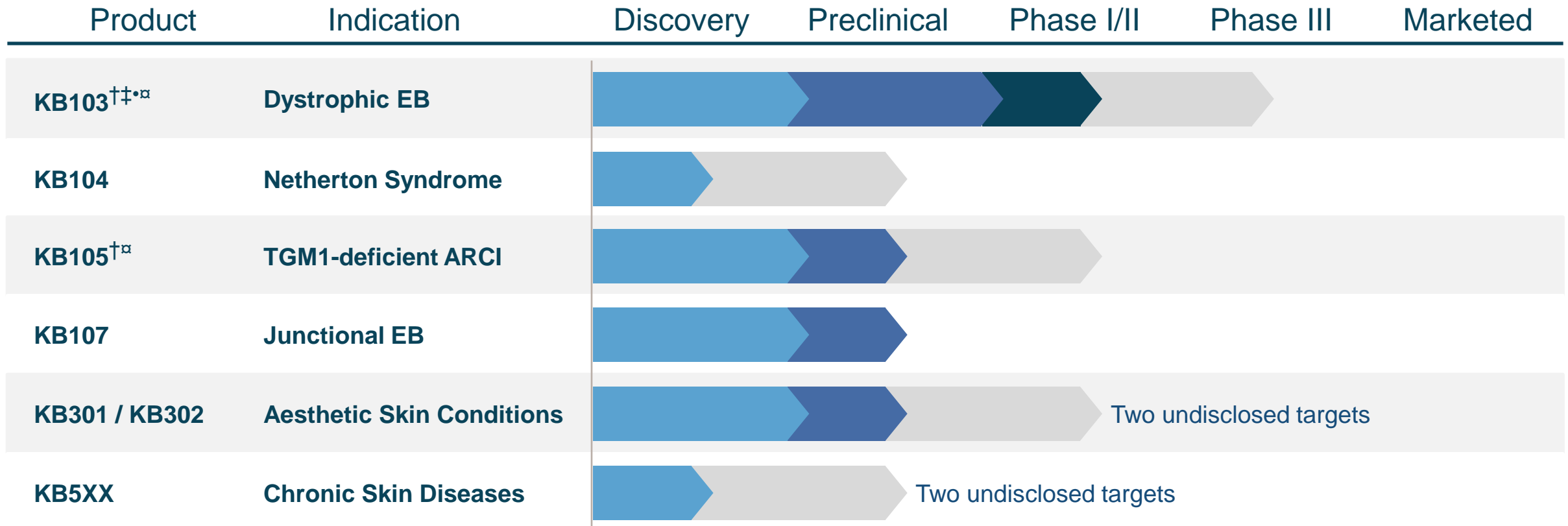
3. Chronic skin diseases

Indications: atopic dermatitis; psoriasis; rosacea; acne



Effectors: Anti-inflammatory antibodies and antibody fragments

Krystal's Current Pipeline



†: FDA Orphan Drug Designation;
 ‡: EMA Orphan Drug Designation;
 •: Fast-track Designation;
 α: FDA Rare Pediatric Disease Designation.



A Catalyst-rich 2019

- Announce final results for KB103 phase I/II trial; 1H 2019
- Commence pivotal phase III trial for KB103; 2H 2019
- Begin phase I clinical trials for two pipeline candidates
 - KB105 for autosomal recessive congenital ichthyosis
 - KB301 for an aesthetic condition
- First GMP facility 'Ancoris' operational.
- Pre-clinical work in pipeline products in anticipation of clinical programs in 2020

A fully-integrated proprietary HSV-1-based gene therapy platform and a pipeline of clinical and non clinical effectors to target skin diseases and conditions

Krystal's Core Competency: CMC/Manufacturing

Established process conducted at Krystal's end-to-end GMP facility

- Maintains control of IP/trade secrets relating to manufacturing process
- Adheres to internal process and production schedules, avoiding use of high demand gene therapy CMOs

Upstream Production Process

- Proprietary engineered vectors and complementary/supporting cell lines developed in-house are used in established methods for production of consistent batches
- Scalable from clinical phase to commercial

Downstream Purification Process

- Work conducted in an aseptic closed system process
- Process accommodates ever-expanding vector pipeline with minimal redevelopment effort between product candidates
- Compliant to global regulatory requirements

KB103*

USAN: *bercolagene telserpavec*

For treatment of dystrophic epidermolysis bullosa (DEB)

- Fast Track Designation Granted
- Orphan Drug Designation in US and EU
- Rare Pediatric Disease Designation in US
- Eligible for Priority Review Voucher
- PRIME Eligibility from EMA

Dystrophic Epidermolysis Bullosa (DEB)

“Butterfly Children” is used to describe young DEB patients because their skin is as fragile as a butterfly’s wings

Dystrophic Epidermolysis Bullosa

A rare, genetic connective tissue disease that causes skin to tear or blister from minor contact

Caused by a mutation in the COL7A1 gene that codes for the COL7 protein

Without COL7 the epidermis does not anchor to the dermis



Epidemiology

Prevalence: Up to 125,000 people are affected by DEB worldwide¹

Incidence: The incidence of DEB is 6.5 per million births in the US²

Current Standard of Care

There are no approved treatments for DEB

Existing therapies limited to expensive and time-consuming palliative treatments

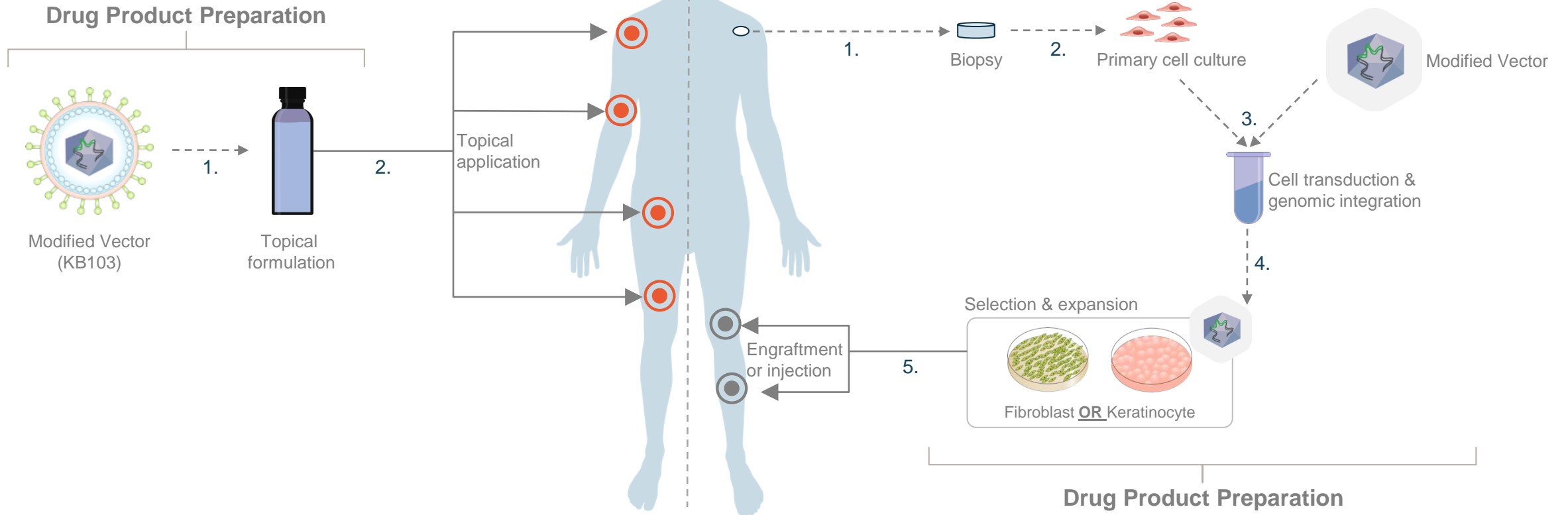
Palliative treatments cost \$200k – \$400k annually^{3,4}

1. DEBRA International, <http://www.debra-international.org/epidermolysis-bullosa/causes-and-subtypes.html>; <http://www.debra-international.org/what-is-eb/causes-and-subtypes/deb.html>
2. Pfendner EG, Lucky AW. Dystrophic Epidermolysis Bullosa. 2006 Aug 21 [Updated 2015 Feb 26]. In: Adam MP, Ardinger HH, Pagon RA, et al., editors. GeneReviews® [Internet].
3. Rashidghamat E., Mellerio J.E., Management of chronic wounds in patients with dystrophic epidermolysis bullosa: challenges and solutions, Chronic Wound Care Management and Research Volume 2017:4, 45-54
4. GENEGRAFT Report Summary. (2015, February 16). Retrieved December 13, 2016, from http://cordis.europa.eu/result/rcn/156078_en.html

Simple, Painless and Easy to Administer



Off-the-shelf, Non-Invasive
Modified HSV-1 Therapy



KB103 Clinical Data

Phase I Trial Design

A Phase I Study of KB103, a Non-Integrating, Replication-Incompetent HSV Vector Expressing the Human Collagen VII Protein, for the Treatment of Dystrophic Epidermolysis Bullosa (DEB)

- Key objectives: Demonstrate efficacy and safety of KB103
- Primary Objectives: Expression of COL7, presence of anchoring fibrils, and safety
- Secondary Objectives: Change in wound area, duration of wound closure, time to wound closure
- Principal investigator: Dr. Peter Marinkovich, MD, Dermatologist, Stanford University
- Trial Design:
 - Randomized, open-label, placebo controlled
 - 2 wounds treated topically: 1 placebo, 1 active
 - 1 intact site treated intradermally
 - Patients were evaluated for COL7 expression by immunofluorescence and for the presence of anchoring fibrils by electron microscopy
 - Initial dosing at Day 0 and a repeat dose a month later; Patient 02 was additionally dosed on Day 14 and Day 42 by PI to understand impact of incremental dose escalation

KB103 Efficacy Update in Wounds With Topical Application

Summary

- Results to date on 2 patients met all primary efficacy (presence of functional COL7 expression as early as Day 2 of treatment, observation of NC1 and NC2 reactive anchoring fibrils and continued expression following repeat administration) and safety endpoints (no adverse events, inflammation or irritation) in topically administered KB103 wounds.
- With respect to secondary endpoints – topically administered KB103 wounds closed in 2 weeks and remained closed through the last timepoint representing 5.7 and 6.6 months of closure, respectively, for Patients 01 and 02. Topically administered placebo treated wounds took 10 weeks to close in Patient 01 and did not completely close throughout the study in Patient 02.
- KB103 treated skin shows presence of functional COL7 expression and anchoring fibrils in both patients.
- Empirical observation that one patient discontinued use of bandages at the site of a KB103-treated area, an area which had required bandages for several months prior to administration.

KB103 Safety Update in Wounds With Topical Application

Summary

KB103 continues to be well tolerated to date following first and repeat dose

- No treatment-related adverse events (serious or otherwise) were reported
- No immune response or blistering observed around the sites of administration following first and repeat dose
- Blood and urine samples collected throughout the study revealed:
 - No systemic viral shedding
 - No adverse events associated with routine labs (chemistry and hematology)
 - No antibodies to COL7 were detected

Phase II Trial Design – Differentiators from Phase I

A Phase II Study of KB103, a Non-Integrating, Replication-Incompetent HSV Vector Expressing the Human Collagen VII Protein, for the Treatment of Dystrophic Epidermolysis Bullosa (DEB)

- Administration to children (vs adults only in Phase 1)
- Administration to larger wounds: up to 20 cm² (vs up to 10 cm² in Phase 1)
- Administration to 2 wounds per patient (vs 1 in Phase 1)
- A higher dose (up to 6 x 10⁸ PFU) is administered per wound
- More frequent administrations allowed, contingent on wound characteristics
- Intradermal administrations were removed

Four patients have been dosed in phase II study

KB105*

For the treatment of Autosomal Recessive
Congenital Ichthyosis associated with TGM1

- Orphan Drug Designation in US
- Rare Pediatric Disease Designation in US
- Eligible for Priority Review Voucher

ARCI Associated With TGM1

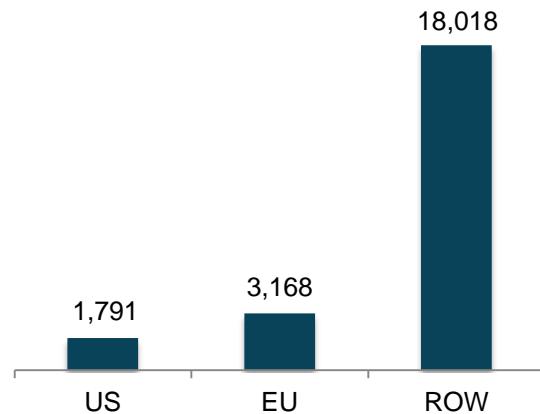
Autosomal Recessive Congenital Ichthyosis (ARCI) associated with TGM1

A condition characterized by thick dry scaly skin, increased trans-epidermal water loss (TEWL), risk for dehydration, sepsis, skin malignancies, etc.

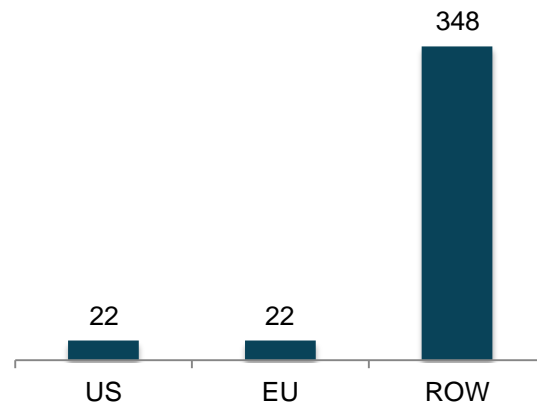
Caused by a mutation of TGM1 gene required for epidermal barrier formation



Prevalence¹⁻⁸



New Cases/Year



Current Standard of Care

There are no approved treatments for ARCI associated with TGM1

Existing approaches limited to time-consuming palliative treatments

1. Rodriguez-Pazos et al. *Actas Dermosifiliogr*. 2013 May;104(4):270–284;
2. Dreyfus et al. *Orphanet J Rare Dis*. 2014 Jan 6;9:1;
3. Hernandez-Martin et al. *J Am Acad Dermatol*. 2012 Aug;67(2):240–244;
4. Pigg et al. *Eur J Hum Genet*. 1998 Nov-Dec;6(6):589–596.

5. Pigg et al. *Acta Derm Venereol*. 2016 Nov 2;96(7):932–937;
6. Orphanet;
7. Foundation for Ichthyosis & Related Skin Types (FIRST);
8. National Organization for Rare Disorders (NORD).

ARCI Associated With TGM1

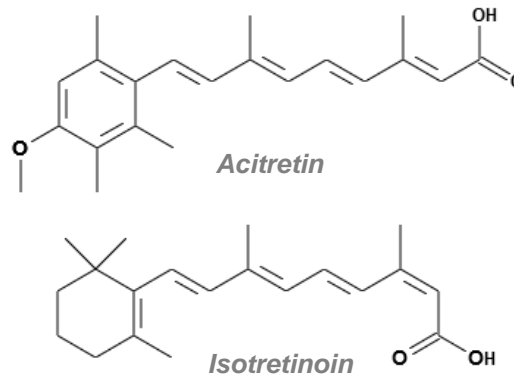
Current standard of care vs. Krystal's approach

Does not address underlying genetic deficiency

Severe adverse effects

- Soft tissue calcification of the joints, e.g., around the spine
- Increased blood triglycerides and cholesterol, potentially inducing or exacerbating atherosclerosis
- Acute and chronic toxicities associated with long-term exposure

Oral retinoids



Particularly ill-suited for certain patient segments

- **Children:** doctors delay retinoid therapy as long as possible due to the bone growth defects (including premature termination of bone elongation) induced by retinoids
- **Women of childbearing age:** retinoids are teratogens (cause fetal abnormalities, miscarriages and severe birth defects) with potentially long half-lives; must be avoided by pregnant women or women who intend on becoming pregnant

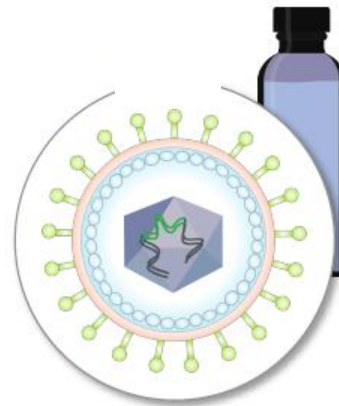
Corrects molecular defects of disease

- KB105 encodes and expresses multiple functional copies of TGM1
- Direct delivery of TGM1 to appropriate skin substrata

Improved safety

- Avoids severe adverse events associated with long-term retinoid therapy

KB105



No systemic exposure to the drug product

Engineered for topical application

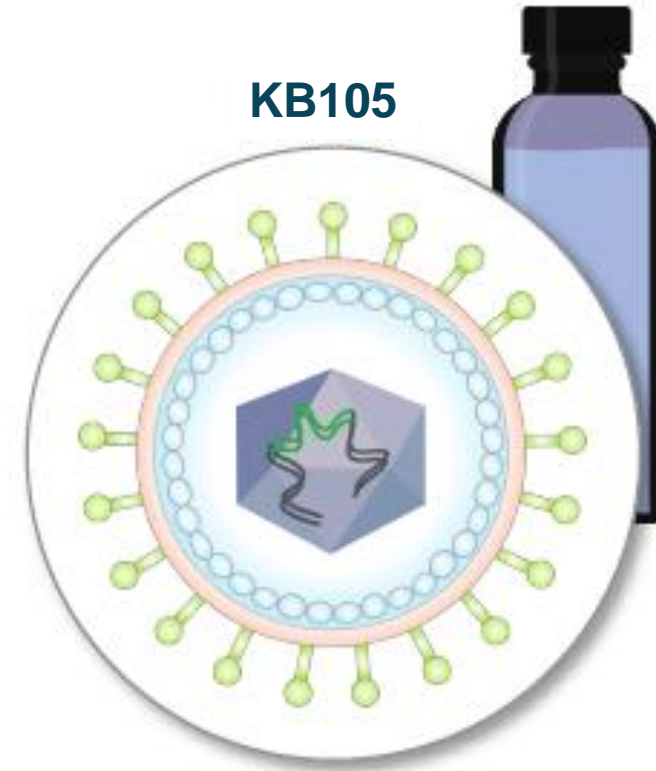
- Can be administered frequently

Suitable for all patient populations

- Including children and women of childbearing age

KB105 Conclusions

- Candidate viruses efficiently transduce target cells and express TGM1 *in vitro* and *in vivo*
- HSV variant shows reduced cytotoxicity
- Human TGM1 consistently expressed at high levels in immunocompetent mice
- KB105 rescues TGM1 expression in ARCI patient cells
- KB105's robust production of TGM1 *in vitro* and *in vivo* supports its use in ARCI patients



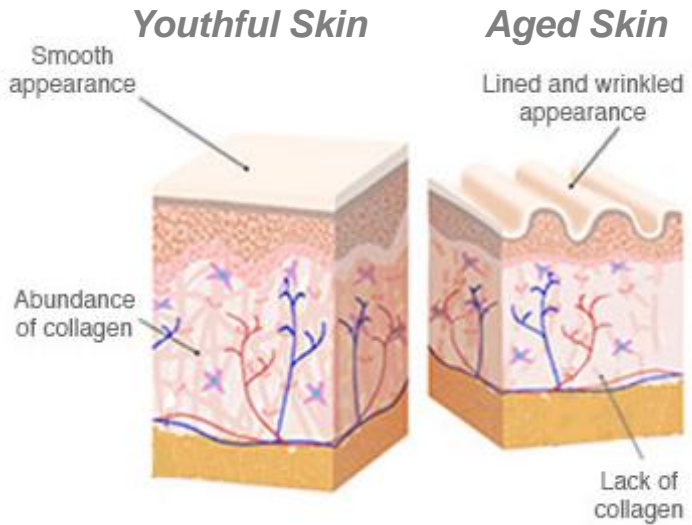
KB105 Timeline to Clinical Readout

Phase I/II clinical trial to begin 2H 2019



Beyond Severe Monogenic Skin Diseases

Application of fully-integrated vector platform to treat aesthetic defects

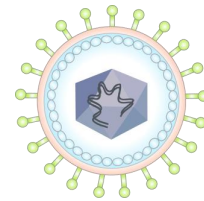


The characteristic features of skin aging are largely due to aberrant collagen homeostasis, resulting in a net collagen deficiency



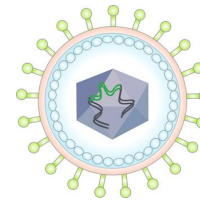
Off-the-shelf, Non-Invasive
Modified HSV-1 Therapy

Vector Platform

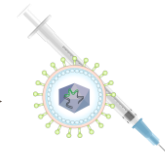


Modify platform via
insertion of human
collagen gene(s)

KB301



Formulate into an
intradermal solution

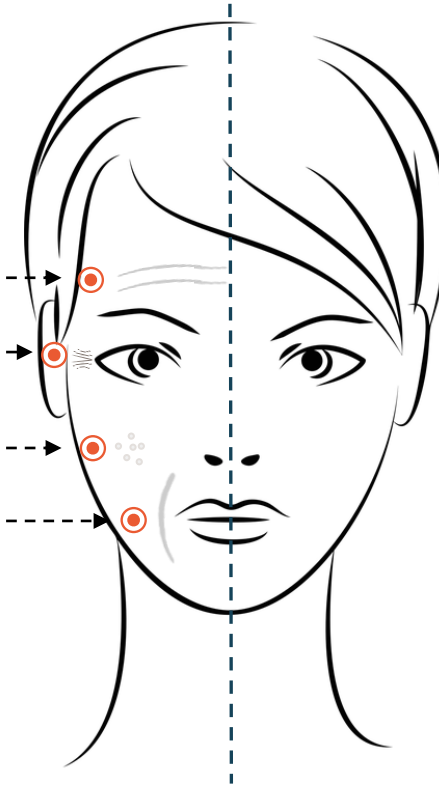


Frown Lines

Crows Feet

Acne Scars

Nasolabial Folds



Fromowitz, J. "Update on Aging Skin"; Florida Society of Dermatology

Beyond Severe Monogenic Skin Diseases

Application of fully-integrated vector platform to treat complex, chronic skin conditions

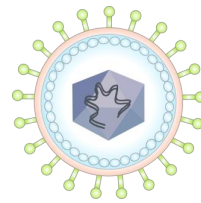
Chronic skin conditions

KB500 series (Antibodies) for Chronic Skin Diseases (Atopic Dermatitis, Psoriasis, etc.)

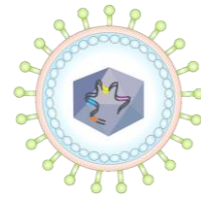


Off-the-shelf, Non-Invasive Modified HSV-1 Therapy

Vector Platform



KB500 series

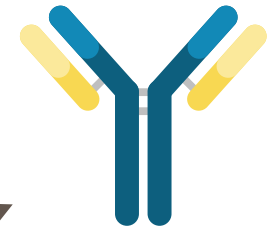


Modify platform via insertion of anti-inflammatory antibody effectors

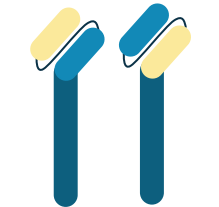
Formulate into a topical gel



Topical



Full-length Abs



Ab Fragments (e.g., scFv-Fcs)



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