



Cautionary Note on Forward-Looking Statements

This presentation contains forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995. All statements other than statements of historical facts are forward-looking statements. In some cases, you can identify forward-looking statements by terms such as "may," "will," "should," "expect," "plan," "anticipate," "could," "future," "outlook," "intend," "target," "project," "contemplate," "believe," "estimate," "predict," "potential," "continue," or the negative of these terms or other similar expressions, although not all forwardlooking statements contain these words. The forward-looking statements in this presentation relate to, among other things, statements regarding the commencement of our planned clinical trials for TWIN, the commencement of our planned bioequivalence study for a generic product candidate, our expected date to report top-line data from our pivotal Phase III clinical programs for Epsolay and TWIN and estimated sales of our product candidates. These statements are neither promises nor guarantees, but involve known and unknown risks, uncertainties and other important factors that may cause our actual results, performance or achievements to be materially different from any future results, performance or achievements expressed or implied by the forward-looking statement, including but not limited to, the following: the fact that we have and expect to continue to incur significant losses; our need for additional funding, which may not be available; our ability to complete the development of our product candidates; our ability to obtain and maintain regulatory approvals for our product candidates in our target markets and the possibility of adverse regulatory or legal actions relating to our product candidates even if regulatory approval is obtained; our ability to commercialize our product candidates; our ability to obtain and maintain adequate protection of our intellectual property; our ability to manufacture our product candidates in commercial quantities, at an adequate quality or at an acceptable cost; our ability to establish adequate sales, marketing and distribution channels; acceptance of our product candidates by healthcare professionals and patients; the possibility that we may face third-party claims of intellectual property infringement; the timing and results of clinical trials that we may conduct or that our competitors and others may conduct relating to our or their products; intense competition in our industry, with competitors having substantially greater financial, technological, research and development, regulatory and clinical, manufacturing, marketing and sales, distribution and personnel resources than we do; potential product liability claims; potential adverse federal, state and local government regulation in the United States, Europe or Israel; and loss or retirement of key executives and research scientists. These and other important factors discussed in the Company's Annual Report on Form 20-F filed with the Securities and Exchange Commission ("SEC") on March 21, 2019 and our other reports filed with the SEC could cause actual results to differ materially from those indicated by the forward-looking statements made in this press release. Any such forward-looking statements represent management's estimates as of the date of this presentation. While we may elect to update such forward-looking statements at some point in the future, unless required by applicable law, we disclaim any obligation to do so, even if subsequent events cause our views to change. Thus, one should not assume that our silence over time means that actual events are bearing out as expressed or implied in such forwardlooking statements. These forward-looking statements should not be relied upon as representing our views as of any date subsequent to the date of this presentation.

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Our Dermatology Company



- Primed to become a global dedicated dermatological company by developing a combination of branded and generic topical drug products
- Expecting results during 2019 from pivotal trials of two branded pipeline candidates,
 based on a proprietary topical microencapsulation delivery system
- Seven established collaborations with two strategic partners on generic candidates already resulted in one approval and one tentative approval by the FDA. First generic product reached the market in February 2019
- Proven track record combined with broad dermatological knowhow

Common Indications Requiring Better Therapies

Acne Vulgaris

- A disease of the pilosebaceous unit, involving abnormalities in sebum production, follicular epithelial desquamation, bacterial proliferation and inflammation
- Benzoyl peroxide (BPO) and tretinoin are mainstay therapies
- Tretinoin is the most widely used Rx topical retinoid, but is rapidly decomposed by BPO and causes irritation
- BPO/tretinoin combination does not currently exist on the market
- ~\$2.7 billion sales in the U.S. in 2018 of several promoted topical brands and many generics, of which fixed-dose combination drugs account for ~\$0.9 billion
- Dermatologists often prefer branded topical drugs even though cheaper generics and OTC alternatives exist

Papulopustular Rosacea

- A chronic, inflammatory skin condition affecting nearly 5 million people in the US
- ~\$0.4 billion sales of topical products in the U.S. in 2018 : Soolantra®, Finacea® and generic metronidazole
- Poor patient adherence to current drugs



Our Branded Drug Product Candidates

TWIN

acne vulgaris

- A cream containing a fixed-dose combination of encapsulated tretinoin and encapsulated benzoyl peroxide
- Major challenges were the instability of tretinoin in the presence of benzoyl peroxide and irritation
- Encapsulation allows stabilization and is also expected to contribute to patient compliance
- Opportunity exists for shift from prescribing tretinoin and existing combinations to prescribing TWIN
- We estimate peak annual sales of \$350M \$400M^(†).

Epsolay®

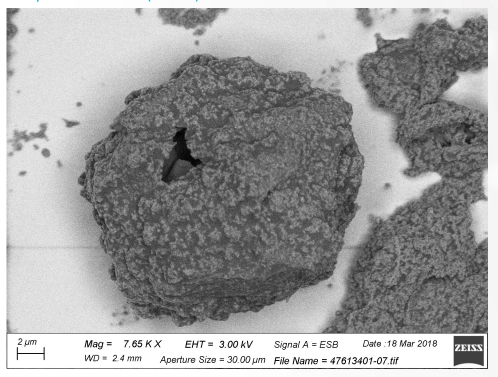
papulopustular rosacea

- A cream containing encapsulated benzoyl peroxide, 5%
- Encapsulation was designed to reduce irritation caused by benzoyl peroxide
- Potential to be the 1st FDA-approved single-active benzoyl peroxide prescription drug product
- We estimate peak annual sales of \$75M \$100M^(†)

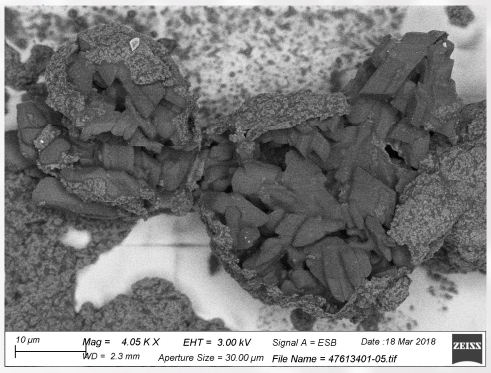


Our Microencapsulation Platform

Encapsulated tretinoin (E-ATRA)



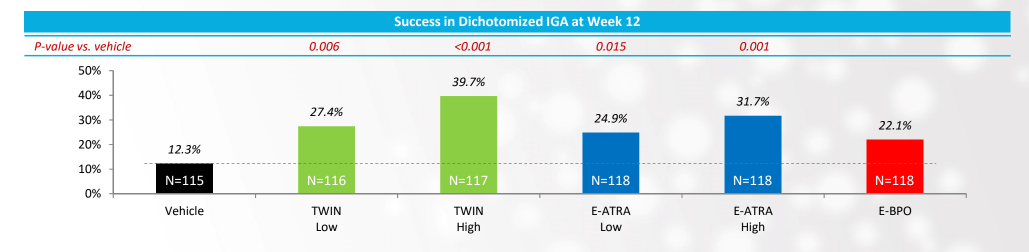
Broken microcapsule containing multiple tretinoin (ATRA) crystals



SEM pictures of our silica-based encapsulated tretinoin

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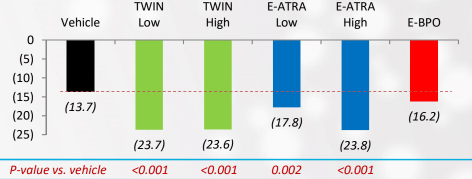
Positive TWIN Factorial Phase II Results (ITT)(†)



Inflammatory Lesion Mean Absolute Change from Baseline at Week 12



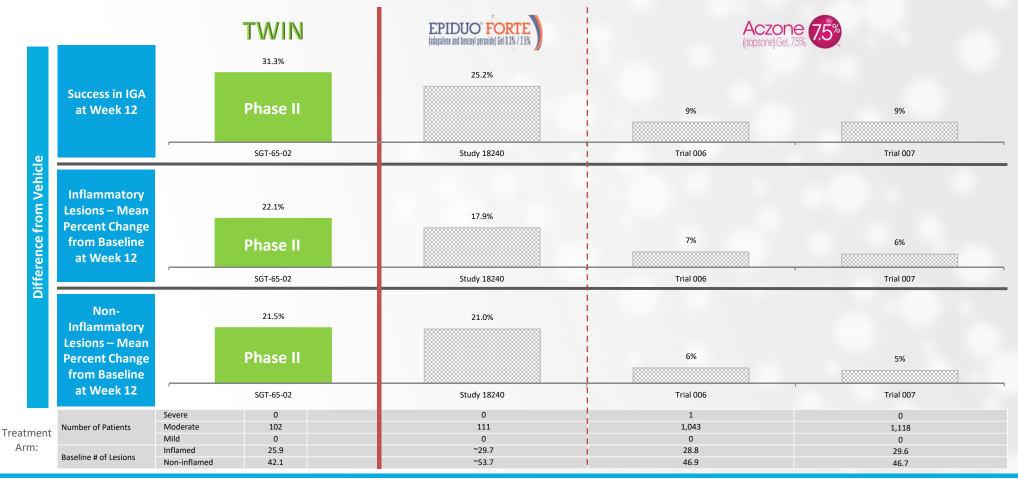
Non-Inflammatory Lesion Mean Absolute Change from Baseline at Week 12





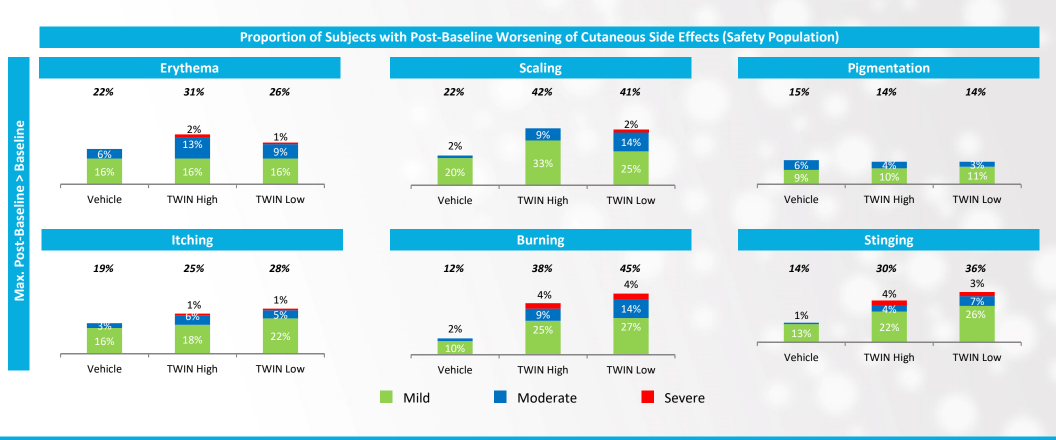
(†) The above calculations were made using Markov Chain Monte Carlo multiple imputation method for handling missing data and without data from one center that discontinued the study. Analyses without imputation (with or without the discontinued center) were highly consistent with the above

Acne Trials Efficacy Results^(†): Moderate Patients





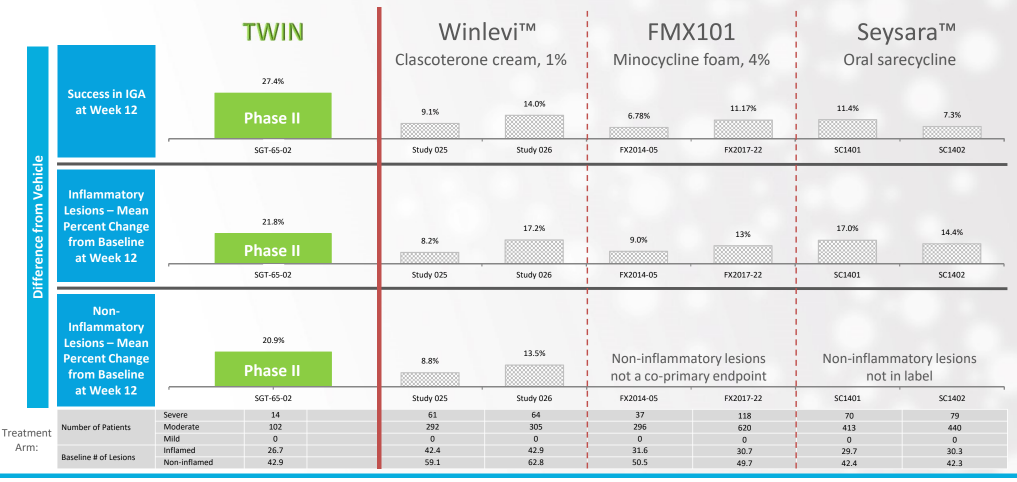
Phase II Cutaneous Tolerability of TWIN





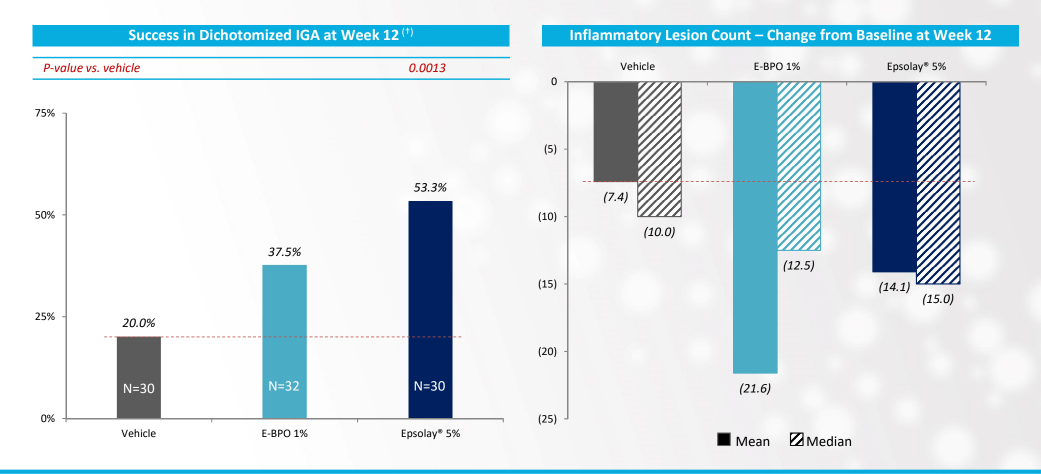
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Efficacy Results of Recent Acne Trials^(†)



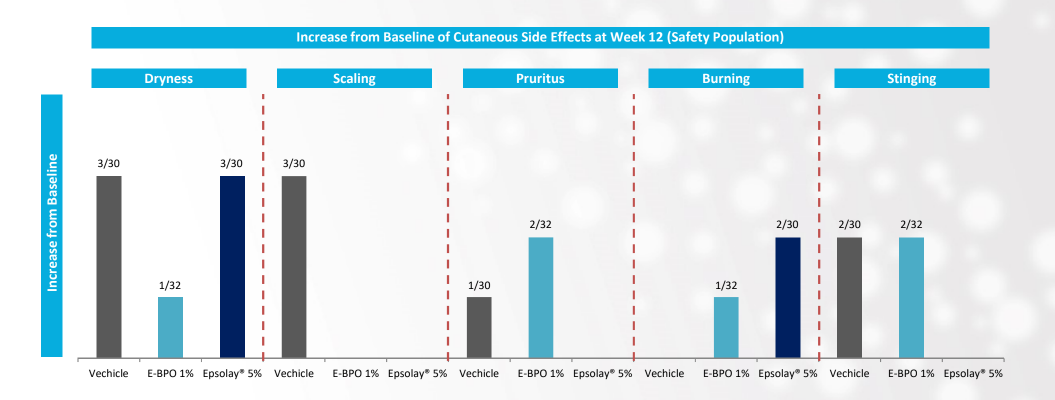


Positive Epsolay® Phase II Results (ITT)





Phase II Cutaneous Tolerability of Epsolay®





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Papulopustular Rosacea Trials Results^(†) (ITT)



(1) "clear" definition: "no inflammatory lesions present with no or very mild erythema immediately localized to and around where inflammatory lesions were present"

(2) 10-week study



Highly-Powered Phase III Trials and Mitigated Risks

TWIN

- Only TWIN and vehicle are required for the pivotal trials, as the requirements of the combination rule act were satisfied in our Phase II trial
- Each pivotal trial is planned to enroll 420 subjects in a 2:1 ratio, with a power of 99%
- No LTSS is required to support our future marketing application, as long as we demonstrate that the systemic exposure of our product is comparable to our reference-listed drug (RLD)
- No pediatric clinical studies are required to support our future marketing application
- Subject to favorable results from our Phase III clinical program, we plan to submit an NDA in 2020

Epsolay®

- Each pivotal trial is planned to enroll 350 subjects in a 2:1 ratio, with a power > 99%
- Long-term safety study (LTSS) was initiated in September 2018
- No pediatric and no Phase I clinical trials are required to support our future marketing application
- Subject to favorable results from our Phase III clinical program, we plan to submit an NDA in 2020

Our Generic Pipeline

Multiple Collaborations

- A portfolio of generic product candidates with favorable commercial agreements that supplement our branded pipeline
- Six collaborations with Perrigo and one with Douglas Pharmaceuticals with 50/50 gross profit sharing

1st Fruitions

- In January 2018, Perrigo received tentative approval from the FDA for ivermectin cream, 1%, developed in collaboration with Sol-Gel. Perrigo was second to file and, as of today, there is no public disclosure of a third filer to the FDA. Sales of RLD reached \$175 million in 2018^(†), and are expected to exceed \$200 million annually by 2020
- In February 2019, Perrigo received approval from the FDA and launched the sale of acyclovir cream, 5%, developed in collaboration with Sol-Gel. As of today, there is no public disclosure of another filer to the FDA. The sales of the RLD were ~\$92 million in 2018^(†)

Recent Developments

Bioequivalence (BE) study results for 5-fluorouracil cream, 5%, expected in 2019

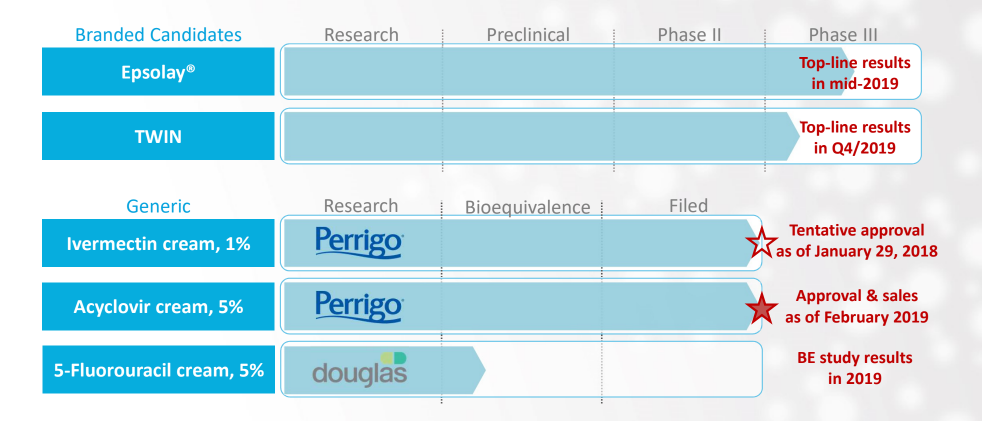


Financial Profile



- Gross proceeds of \$86.3 million raised in IPO of 7,187,500 ordinary shares on February 5, 2018
- 18,949,968 shares outstanding as of December 31, 2018
- \$63.0 million of cash and investments as of December 31, 2018
- Cash runway expected to be sufficient to fund Phase III clinical programs for TWIN and Epsolay[®], a bioequivalence study, and our activities until the end of Q2/2020

Our Upcoming Milestones





Recent Milestones and Next Steps

2017 2018 2019 Reported positive results from TWIN Phase II Obtained tentative ANDA approval Obtained ANDA approval trial in acne vulgaris for ivermectin cream (sponsored by Perrigo) for acyclovir cream (sponsored by Perrigo) Had an EoPII meeting with the FDA about Had an EoPII meeting with the FDA about Launched acyclovir cream (by Perrigo) Epsolay® TWIN and addressed the combination rule act Submitted a Paragraph IV ANDA, for Initiated Epsolay® Phase III program Plans to report Phase III results for Epsolay® ivermectin cream, 1% (sponsored by Perrigo) in papulopustular rosacea in papulopustular rosacea Plans to report Phase III results for TWIN Initiated LTSS for Epsolay® in acne vulgaris Initiated TWIN Phase III program Plans to report BE study results for 5-fluorouracil cream, 5% in acne vulgaris Initiated a bioequivalence study for 5-fluorouracil cream, 5% in actinic keratosis Hired U.S. commercialization leader for the launches of TWIN and Epsolay®





NASDAQ: SLGL

www.sol-gel.com