
UNITED STATES
SECURITIES AND EXCHANGE COMMISSION
Washington, D.C. 20549

FORM 6-K

Report of Foreign Private Issuer
Pursuant to Rule 13a-16 or 15d-16
Under the Securities Exchange Act of 1934

For the month of June 2021

Commission File Number 001-38367

SOL-GEL TECHNOLOGIES LTD.

(Translation of registrant's name into English)

7 Golda Meir Street
Ness Ziona 7403650, Israel
(Address of principal executive offices)

Indicate by check mark whether the registrant files or will file annual reports under cover Form 20-F or Form 40-F.

Form 20-F Form 40-F

Indicate by check mark if the registrant is submitting the Form 6-K in paper as permitted by Regulation S-T Rule 101(b)(1):

Indicate by check mark if the registrant is submitting the Form 6-K in paper as permitted by Regulation S-T Rule 101(b)(7):

INFORMATION CONTAINED IN THIS REPORT ON FORM 6-K

Sol-Gel Technologies Ltd. (the "Company") is posting on its website a corporate presentation.

Attached hereto and incorporated by reference in this Report on Form 6-K is the following exhibit:

[Exhibit 99.1: Corporate Presentation.](#)

SIGNATURES

Pursuant to the requirements of the Securities Exchange Act of 1934, the Registrant has duly caused this report to be signed on its behalf by the undersigned, thereunto duly authorized.

SOL-GEL TECHNOLOGIES LTD.

Date: June 22, 2021

By: /s/ Gilad Mamlok
Gilad Mamlok
Chief Financial Officer



Sol-Gel

Advanced Topical Therapy

NASDAQ: SLGL

June 2021



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 forward-looking statements contain these words. The forward-looking statements in this presentation relate to, among other things, statements regarding the PDUFA goal date for TWYNEO, approval and commercial launch of EPSOLAY and TWYNEO, and the negotiations with a potential partner regarding the commercialization of EPSOLAY and TWYNEO, anticipated timing of results of the ongoing Phase 1 clinical trial of SGT-210, the expectation to launch a partnered generic drug starting in the second quarter of 2021, our expectations regarding our liquidity and ability to fund operational and capital expenditure requirements, 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: risks relating to the timing of the PDUFA action date for EPSOLAY; the timing of FDA approval, if any, of EPSOLAY and TWYNEO; the risk that we may not execute an agreement for the commercialization of EPSOLAY and TWYNEO and risks related to the terms thereof; 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, and obtain marketing approval for, 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 and launch our product candidates at all or on a timely basis; 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; delays in the launch of product candidates and generic drugs; intense competition in our industry; potential product liability claims; potential adverse federal, state, and local government regulation in the United States, Europe, or Israel; the impact of pandemics, such as COVID-19 (coronavirus); 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 4, 2021, and in 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 presentation. 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 forward-looking 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. This presentation contains trademarks, trade names, and service marks of other companies, which are the property of their respective owners. We do not intend our use or display of other parties’ trademarks, trade names, or service marks to imply, and such use or display should not be construed to imply, a relationship with, or endorsement or sponsorship of us by, these other parties.

TECHNOLOGY

- Proprietary silica-based microencapsulation technology

EPSOLAY®

- PDUFA goal date was set for April 26, 2021. Awaiting FDA's pre-approval inspection
- Potential to be the first single-active BPO approved by the FDA as a prescription drug product

TWYNEO®

- PDUFA goal date set for August 1, 2021
- Potential to be first FDA-approved acne treatment that contains fixed-dose combination of BPO and tretinoin

SGT-210

- Phase I proof-of-concept study for erlotinib gel in palmoplantar keratoderma was completed

EARLY STAGE

- Pending patent applications for tapinarof and roflumilast in various skin conditions

GENERIC

- Twelve 50/50 gross profit-sharing collaborations with Perrigo
- \$0.7 million in net revenues in 1Q/21



THE SCIENCE BEHIND OUR PROPRIETARY TECHNOLOGY

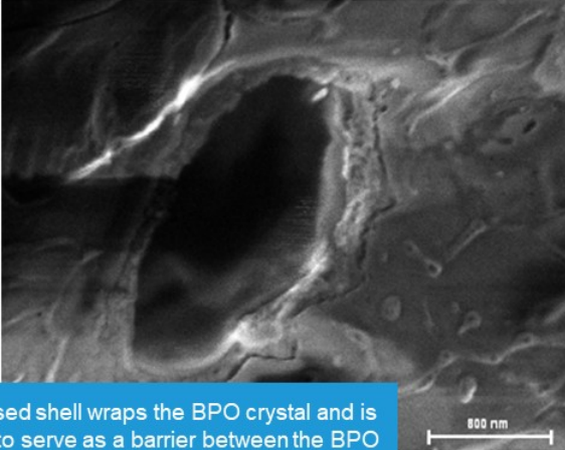
Aiming to provide effective and tolerable topical
therapies to achieve local action



ENCAPSULATION IS DESIGNED TO ALLOW FOR CONTINUOUS FLOW

ENCAPSULATED BENZOYL PEROXIDE (E-BPO)

CRYO-SEM PICTURE



Silica-based shell wraps the BPO crystal and is intended to serve as a barrier between the BPO crystals and the skin

ENERGY-DISPERSIVE X-RAY SPECTROSCOPY MAPPING

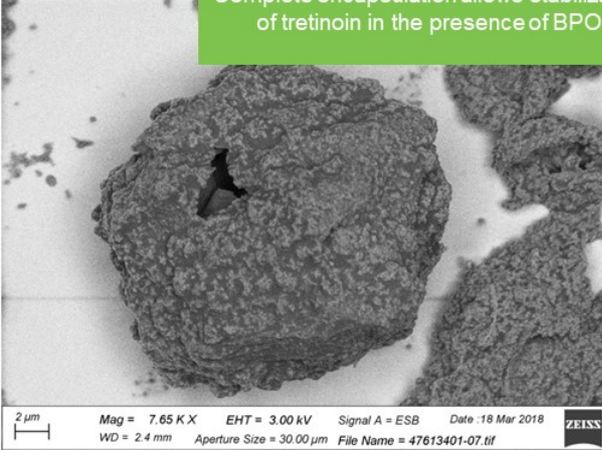


After application onto skin, BPO slowly migrates through the shell resulting in a continuous flow of BPO for up to 24 hours

ENCAPSULATION IS DESIGNED TO ENHANCE STABILITY
ENCAPSULATED TRETINOIN (E-TRETINOIN)

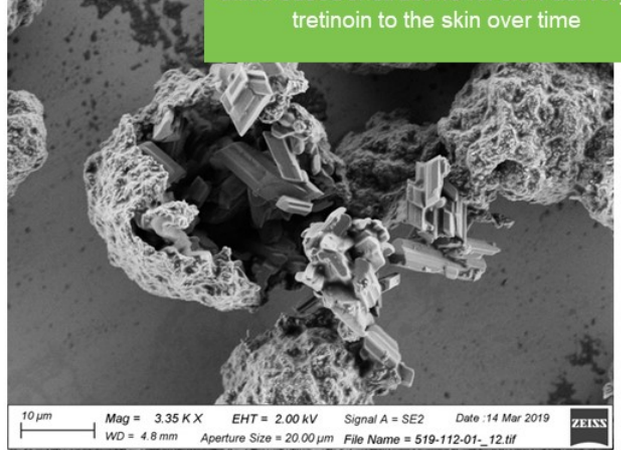
SEM PICTURE

Complete encapsulation allows stabilization of tretinoin in the presence of BPO



SEM PICTURE

Silica-based shell allows for slow delivery of tretinoin to the skin over time





THE CHALLENGE

**CHRONIC
CONDITION WITH
POOR ADHERENCE
TO CURRENT
TREATMENTS**

**UNMET NEED IN
PAPULOPUSTULAR
ROSACEA**



Papulopustular Rosacea

Chronic, inflammatory condition that primarily affects the face and is often characterized by flushing, redness, inflamed bumps, and pustules

How is it Treated?

- Topical antimicrobials (metronidazole, clindamycin)
- Topical anti-mite (ivermectin)
- Systemic antibiotics (minocycline, doxycycline)

Current Treatment Shortfalls

- Insufficient efficacy resulting in poor adherence
- Systemic side effects
- Contributing to antibiotic resistance

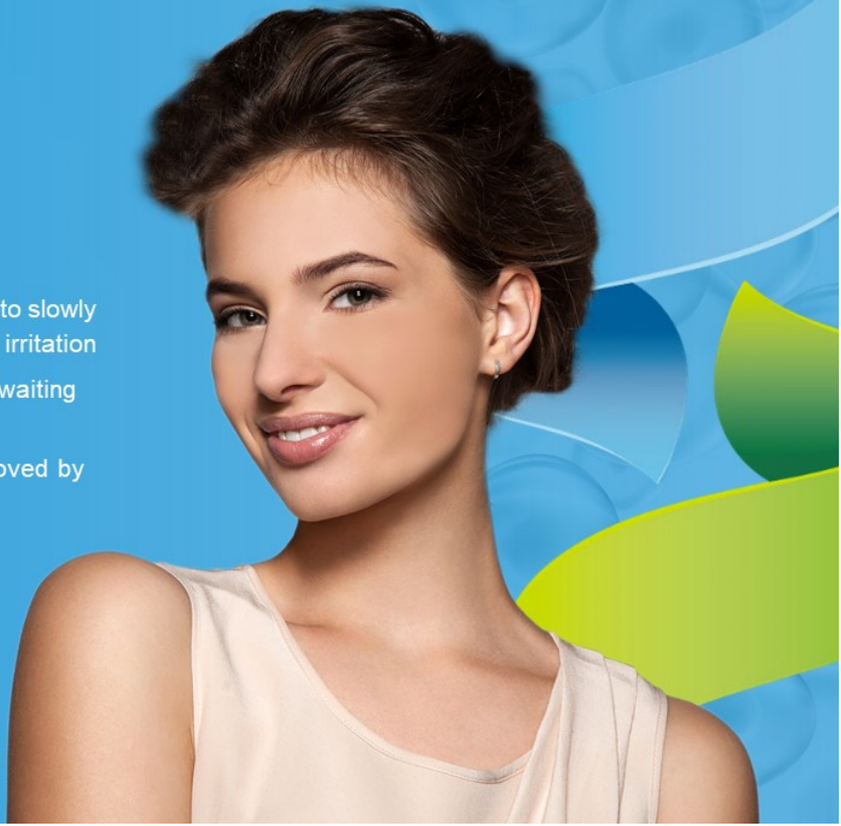
SOL-GEL SOLUTION*

EPSOLAY[®]

Benzoyl Peroxide Cream, 5%

- Encapsulation was designed to allow the BPO to slowly migrate from the microcapsules to help reduce irritation
- PDUFA goal date was set for April 26, 2021. Awaiting FDA's pre-approval inspection
- Potential to be the first single-active BPO approved by the FDA as a prescription drug product

* EPSOLAY is investigational. Safety and efficacy have not been established





EPSOLAY[®]
PHASE III STUDIES

Two Parallel, Multicenter, Double-Blinded,
Randomized, Vehicle-Controlled Studies, 2:1 Ratio, QD



PHASE III DESIGN

TWO CO-PRIMARY EFFICACY ENDPOINTS AT WEEK 12

Inclusion Criteria	≥18 years old; "Moderate" or "Severe" rosacea; ≥15 to ≤70 inflammatory lesions; ≤2 nodules
How is it Treated?	Weeks 2, 4, 8, 12 (end of study)
Investigator Global Assessment (IGA) Definition	<ul style="list-style-type: none">• "Clear": Skin clear of inflammatory papules or pustules• "Almost Clear": Very few small papules or pustules and very mild dull erythema is present• "Mild": Few small papules or pustules and mild dull or light pink erythema is present• "Moderate": Several to many small or larger papules or pustules and moderate light to bright red erythema is present• "Severe": Numerous small and/or larger papules or pustules and severe erythema that is bright red to deep red is present
Primary Endpoints	<ul style="list-style-type: none">• Proportion of patients with IGA "Clear" or "Almost Clear" relative to baseline at Week 12• Absolute mean change in inflammatory lesion counts from baseline to Week 12



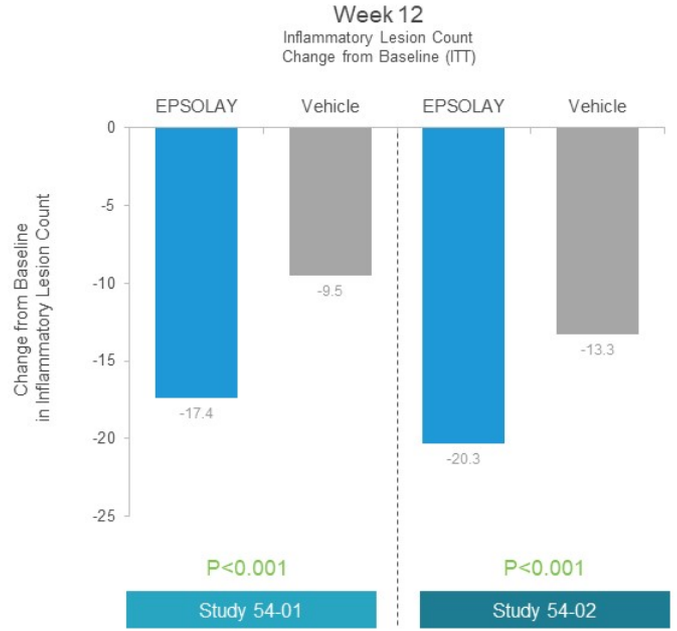
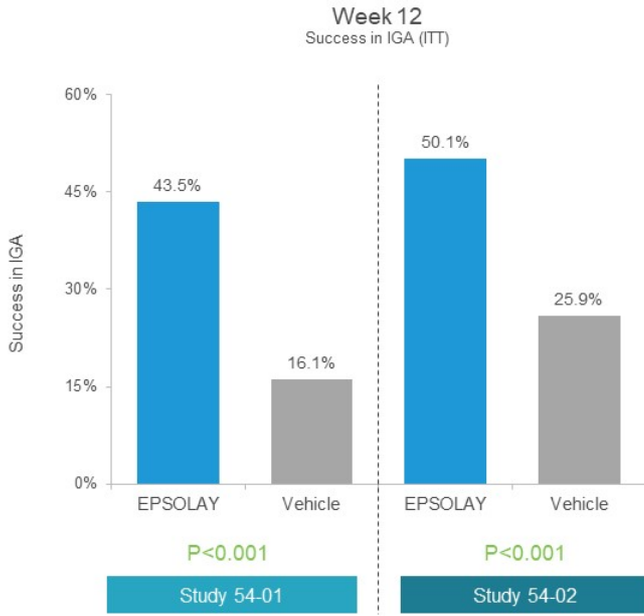
WELL-BALANCED CLINICAL STUDIES

Baseline, Discontinuation & Completion		Study 54-01		Study 54-02	
		EPSOLAY	Vehicle	EPSOLAY	Vehicle
Baseline	IGA "Moderate" Subjects	210 (86.4%)	104 (88.1%)	227 (90.8%)	112 (91.8%)
	IGA "Severe" Subjects	33 (13.6%)	14 (11.9%)	23 (9.2%)	10 (8.2%)
	Mean Inflammatory Lesion Count (SD)	25.7 (11.07)	26.3 (12.45)	29.8 (14.00)	27.5 (13.04)
	Median Inflammatory Lesion Count (range)	22.0 (15-69)	21.0 (15-70)	25.0 (15-70)	22.5 (15-70)
Discontinued Subjects	Withdrawal by Subject	9	3	9	4
	Adverse Events	5	1	4	0
	Lost to Follow-Up	6	6	1	4
	Pregnancy/Protocol Violation/Other	1	1	1	1
Intention-to-Treat (ITT)		243	118	250	122

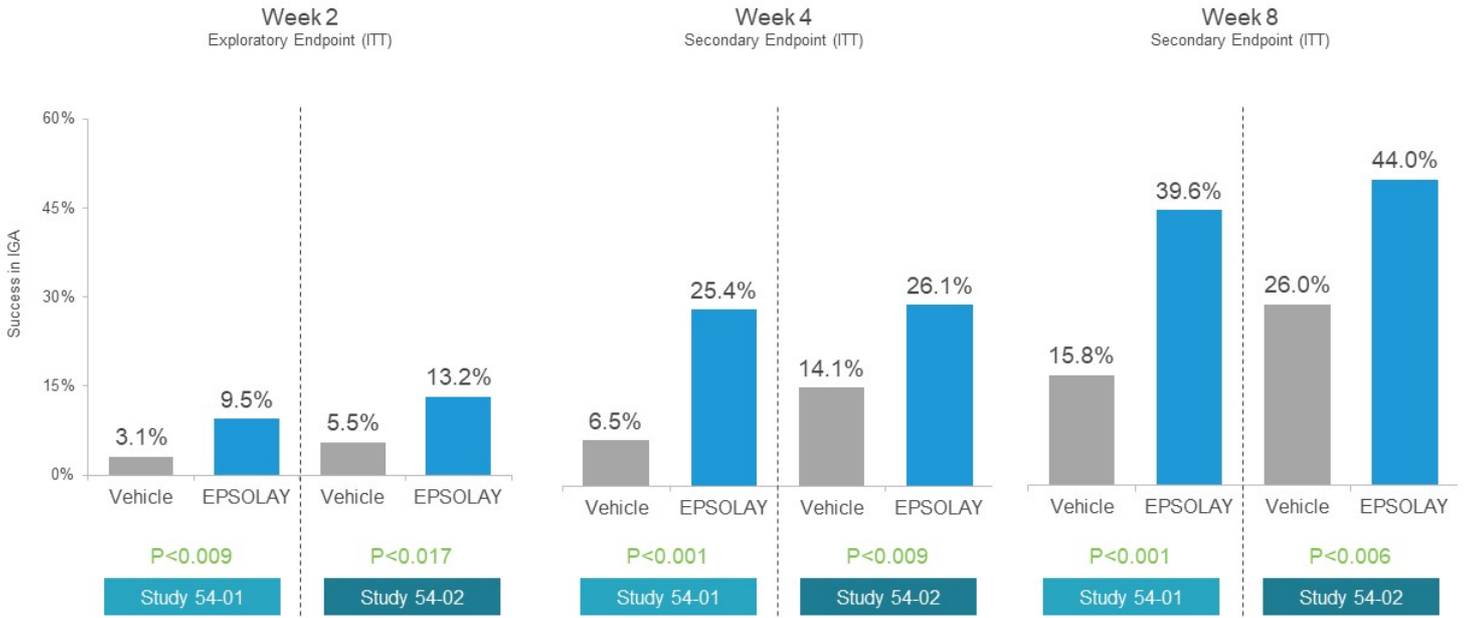
SD = Standard Deviation



SUCCESS IN PRIMARY ENDPOINTS



SUCCESS IN IGA
IMPROVEMENT AS OF WEEK 2





IMPROVEMENT AS OF WEEK 2



Subject 116-009 || 41 years old | Female | White | Not Hispanic or Latino*

ONSET OF ACTION AS OF WEEK 2

BASELINE



"Severe"; 31 inflamed lesions

WEEK 2



"Clear"; No inflamed lesions

WEEK 4



"Clear"; No inflamed lesions

WEEK 8



"Clear"; No inflamed lesions

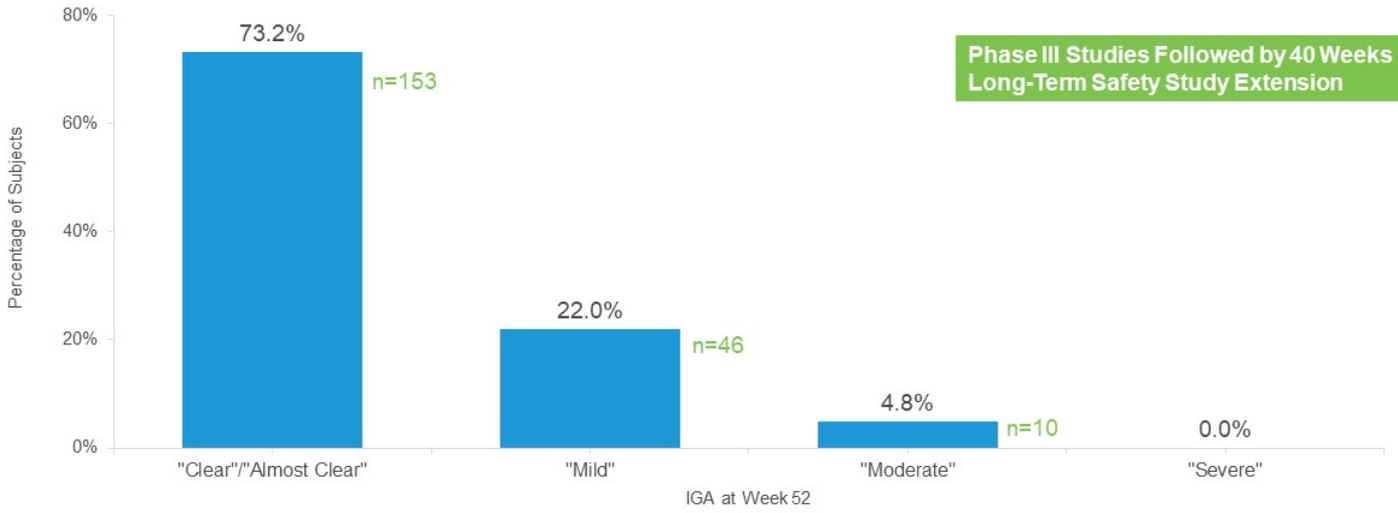
WEEK 12



"Almost Clear"; 1 inflamed lesion

* Individual results vary

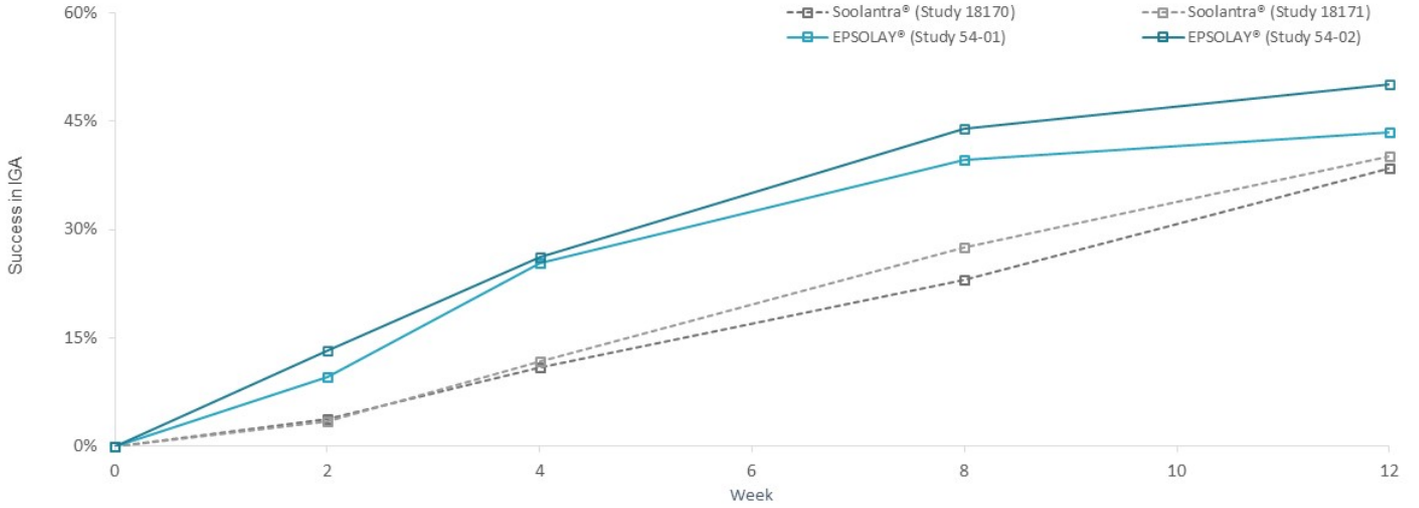
LONG-TERMSAFETY STUDY
IMPROVEMENT IN IGA*



* This study was not designed for efficacy; however, efficacy was evaluated. Interpret results with caution

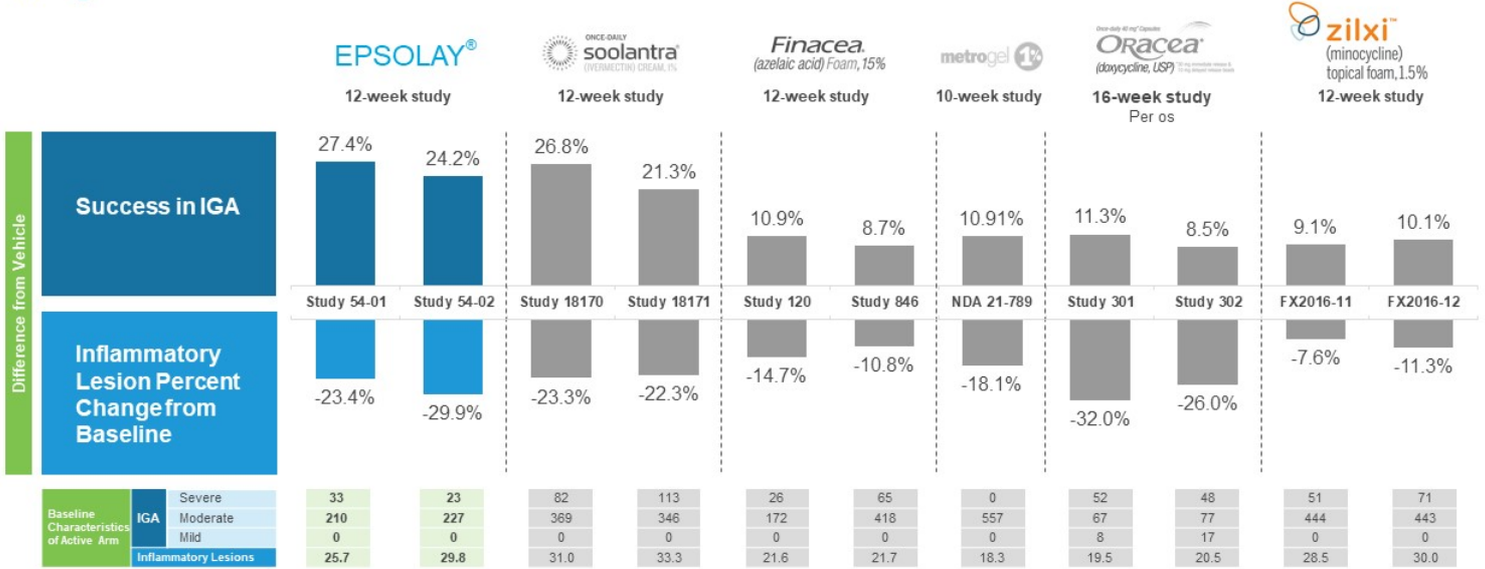


SIDE-BY-SIDE WITH HISTORICAL RESULTS* IMPROVEMENT OVER TIME



* Sol-Gel did not conduct a head-to-head comparison trial or study. The results described above are for illustrative purposes only and should not be construed as conclusions to be drawn as if we conducted a head-to-head comparison trial or study

SIDE-BY-SIDE WITH HISTORICAL RESULTS*
PRIMARY ENDPOINTS



* Sol-Gel did not conduct a head-to-head comparison trial or study. The results described above are for illustrative purposes only and should not be construed as conclusions to be drawn as if we conducted a head-to-head comparison trial or study



PRIMARILY MILD-TO-MODERATE

TREATMENT-EMERGENT ADVERSE EVENTS

Subjects with Treatment-Emergent Adverse Events (TEAEs)	Study 54-01		Study 54-02	
	EPSOLAY (n=239)	Vehicle (n=113)	EPSOLAY (n=249)	Vehicle (n=120)
Treatment-Related Mild & Moderate TEAEs	12 (5%) [^]	3 (2.7%) [^]	8 (3.2%) [^]	0
Treatment-Related Severe TEAEs	2 (0.8%) [≠]	0	1 (0.4%) [*]	0
Not-Related TEAEs	35 (14.6%)	14 (12.4%)	41 (16.5%)	22 (18.2%)
Not-Related Serious TEAEs	0	1 (0.9%) [†]	1 (0.4%) [‡]	0

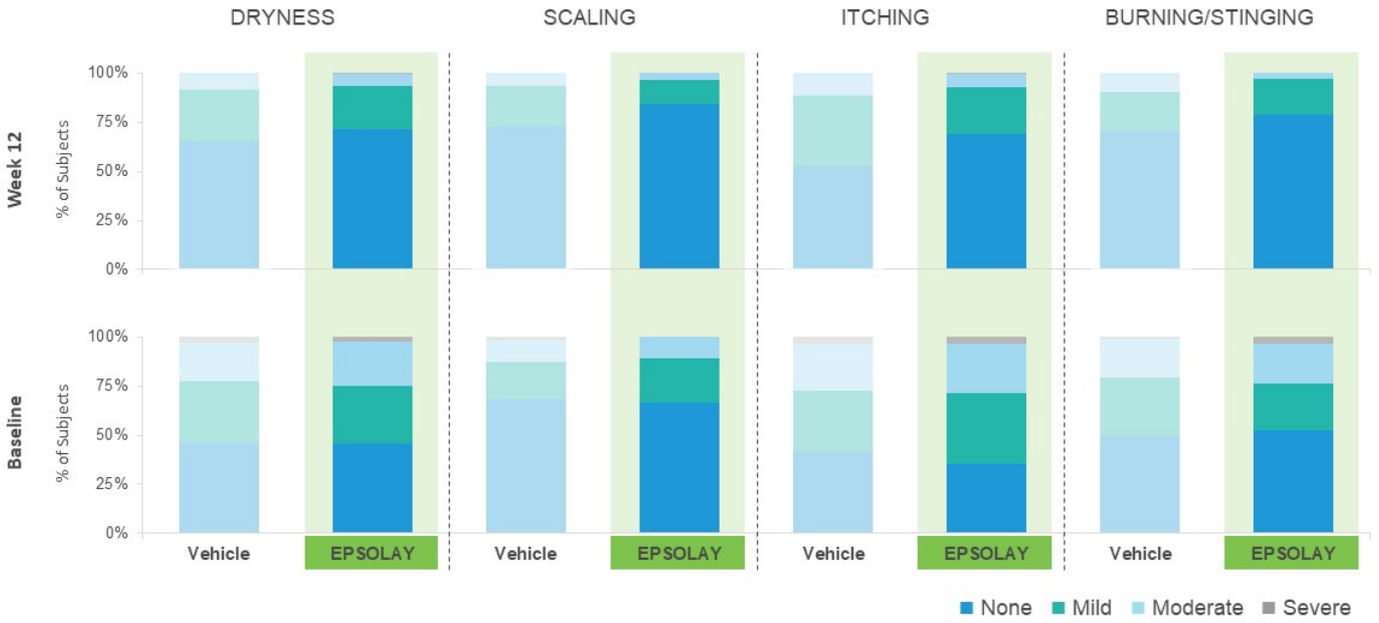
[^] Most frequently reported adverse events being application site erythema, pain and pruritus

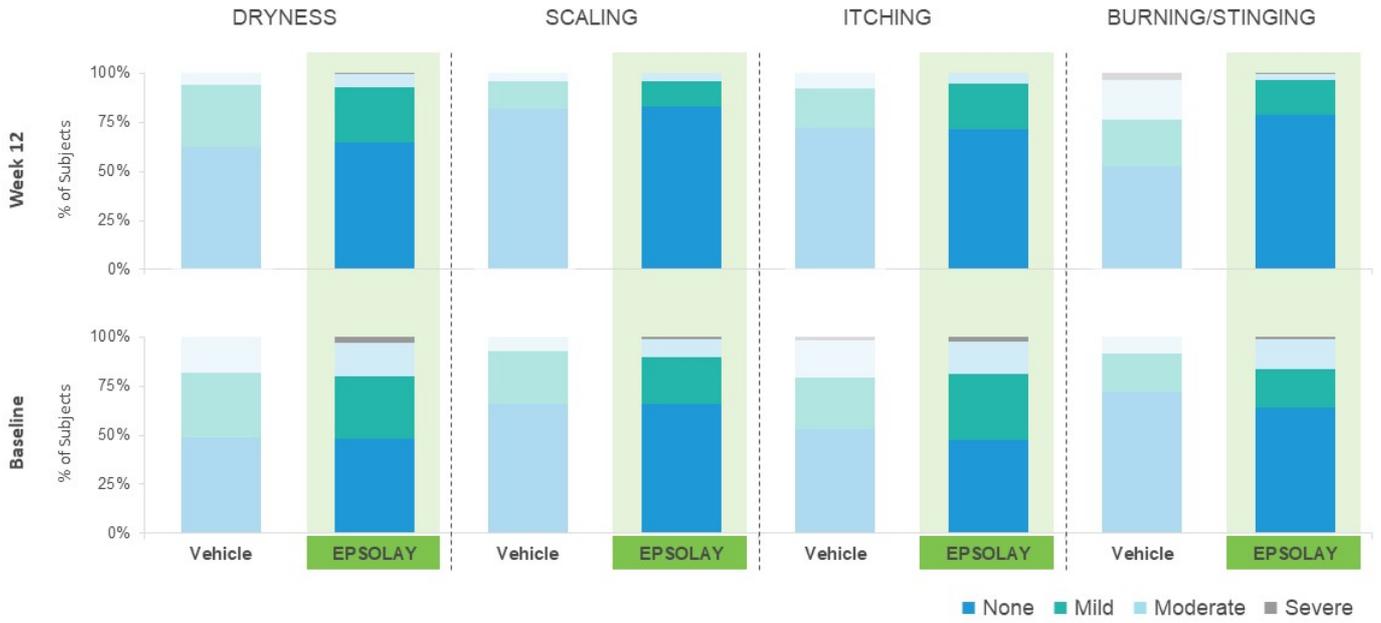
[≠] One subject with application site erythema and another with application site pruritus and pain

^{*} One subject with application site erythema

[†] One subject with femur fracture

[‡] One subject with spinal compression fracture







THE CHALLENGE

MULTIFACTORIAL
DISEASE
REQUIRING
POWERFUL
COMBINATION
TREATMENTS

UNMET NEED IN
ACNE VULGARIS



Acne Vulgaris

A multifactorial disease of the pilosebaceous unit, involving abnormalities in sebum production, follicular epithelial desquamation, bacterial proliferation, and inflammation

How is it Treated?

- Topical BPO, retinoids (such as tretinoin, adapalene), antibiotics, and their combinations
- Oral Isotretinoin and antibiotics

Current Treatment Shortfalls

- Insufficient efficacy negatively affects self-esteem
- Systemic side effects
- Contributes to antibiotic resistance

SOL-GEL SOLUTION*

TWYNEO[®]

Benzoyl Peroxide 3% & Tretinoin 0.1%, Cream

- Encapsulation was designed to stabilize tretinoin and to enable both tretinoin and BPO to slowly migrate from their microcapsules to help reduce irritation
- PDUFA goal date set for August 1, 2021
- Potential to be first FDA-approved acne treatment that contains fixed-dose combination of BPO and tretinoin

* TWYNEO is investigational. Safety and efficacy have not been established





TWYNEO[®]
PHASE III STUDIES

Two Parallel, Multicenter, Double-Blinded,
Randomized, Vehicle-Controlled Studies, 2:1 Ratio, QD



THREE CO-PRIMARY EFFICACY ENDPOINTS AT WEEK 12

Inclusion Criteria	≥9 years old; "Moderate" or "Severe" acne; ≥20 to ≤100 inflammatory lesions; ≥30 to ≤150 non-inflammatory lesions; ≤2 cysts/nodules
Visits	Weeks 2, 4, 8, 12 (end of study)
Investigator Global Assessment (IGA) Definition	<ul style="list-style-type: none">• "Clear": Normal, clear skin with no evidence of acne vulgaris• "Almost Clear": Rare non-inflammatory lesions present, with rare non-inflamed papules (papules must be resolving and may be hyperpigmented, though not pink-red)• "Mild": Some non-inflammatory lesions are present, with few inflammatory lesions (papules/pustules only; no nodulo-cystic lesions)• "Moderate": Multiple Non-inflammatory lesions and, inflammatory lesions are evident (several to many comedones and papules/pustules, and there may or may not be one small nodulo-cystic lesion)• "Severe": Inflammatory lesions are more apparent, many comedones and papules/pustules, there may or may not be a few nodulo-cystic lesions
Primary Endpoints	<ul style="list-style-type: none">• Proportion of subjects with an assessment of "Clear" or "Almost Clear" and with at least a 2-grade improvement in IGA from baseline at Week 12• Absolute change in inflammatory lesion counts from baseline at Week 12• Absolute change in non-inflammatory lesion counts from baseline at Week 12

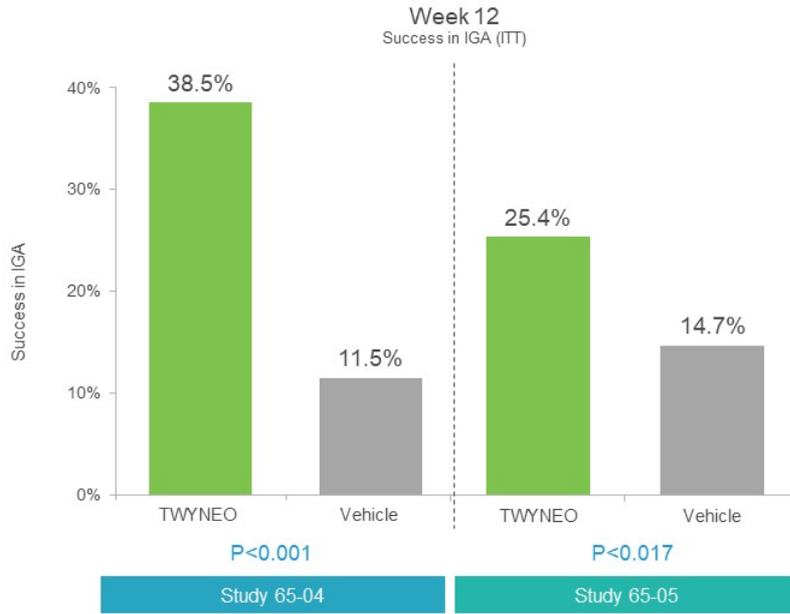


WELL-BALANCED CLINICAL STUDIES

Baseline, Discontinuation & Completion		Study 65-04		Study 65-05	
		TWYNEO	Vehicle	TWYNEO	Vehicle
Baseline	IGA "Moderate" Subjects	251 (89.3%)	132 (92.3%)	262 (90.3%)	133 (93.0%)
	IGA "Severe" Subjects	30 (10.7%)	11 (7.7%)	28 (9.7%)	10 (7.0%)
	Mean Inflammatory Lesion Count (SD)	33.5 (14.62)	33.5 (14.69)	28.2 (8.70)	27.5 (8.52)
	Median Inflammatory Lesion Count (range)	28.0 (20-92)	28.0 (20-90)	25.0 (20-62)	25 (20-75)
	Mean Non-Inflammatory Lesion Count (SD)	48.6 (20.24)	47.1 (19.97)	44.6 (18.03)	44.9 (18.82)
	Median Non-Inflammatory Lesion Count (range)	42.0 (30-148)	41.0 (30-140)	39.0 (23-149)	38.0 (30-123)
Discontinued Subjects	Withdrawal by Subject/Parent/Guardien	13	5	18	5
	Adverse Events	4	0	12	0
	Lost to Follow-Up	10	7	15	7
	Pregnancy/Protocol Violation/Physician Decision/Other	5	0	3	0
Intention-to-Treat (ITT)		281	143	290	144

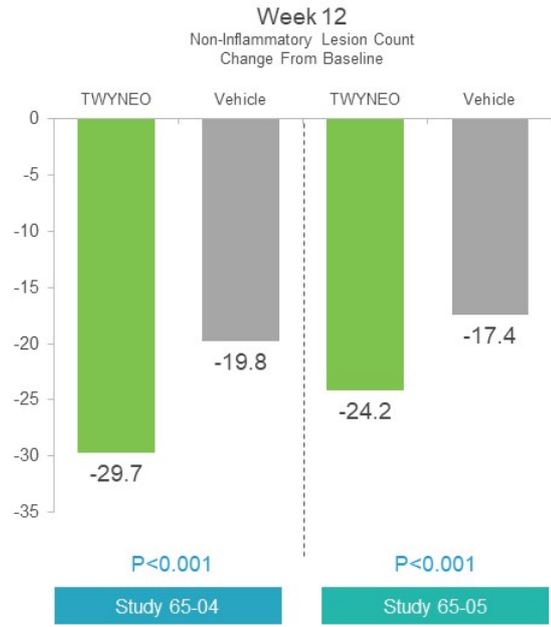
SD = Standard Deviation

PHASE III RESULTS
SUCCESS IN IGA





SUCCESS IN REDUCING LESIONS



Subject 507-003 || 18 years old | Female | White | Not Hispanic or Latino*

IMPROVEMENT IN SEVERE PATIENT

BASELINE



"Severe": 29 inflamed lesions
31 non-inflamed lesions; 1 nodule

WEEK 12

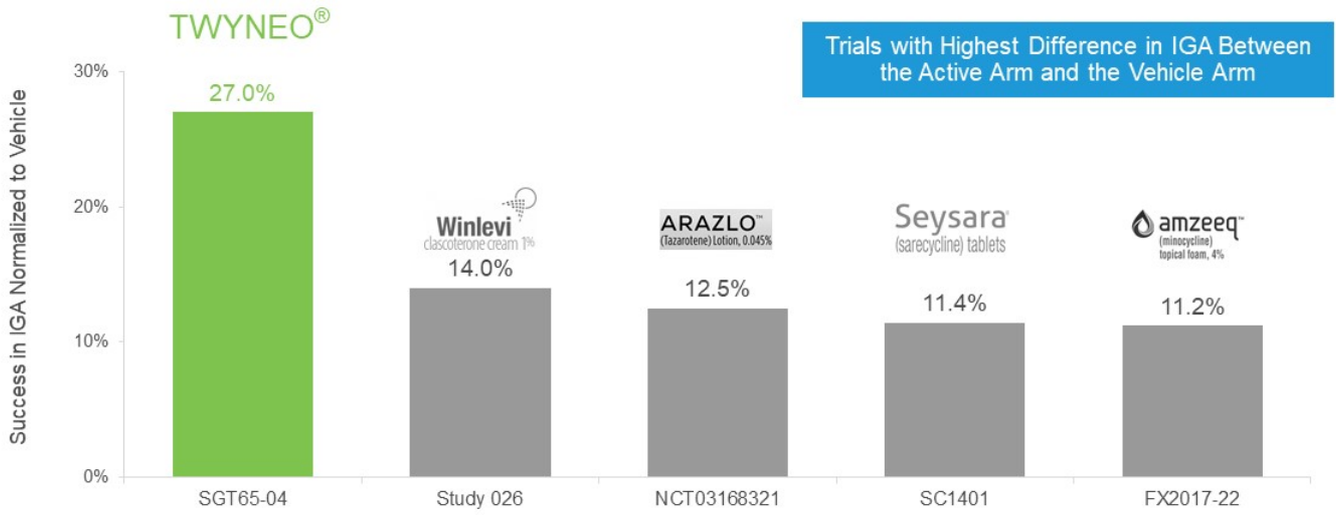


"Moderate": 9 inflamed lesions
5 non-inflamed lesions; No nodules

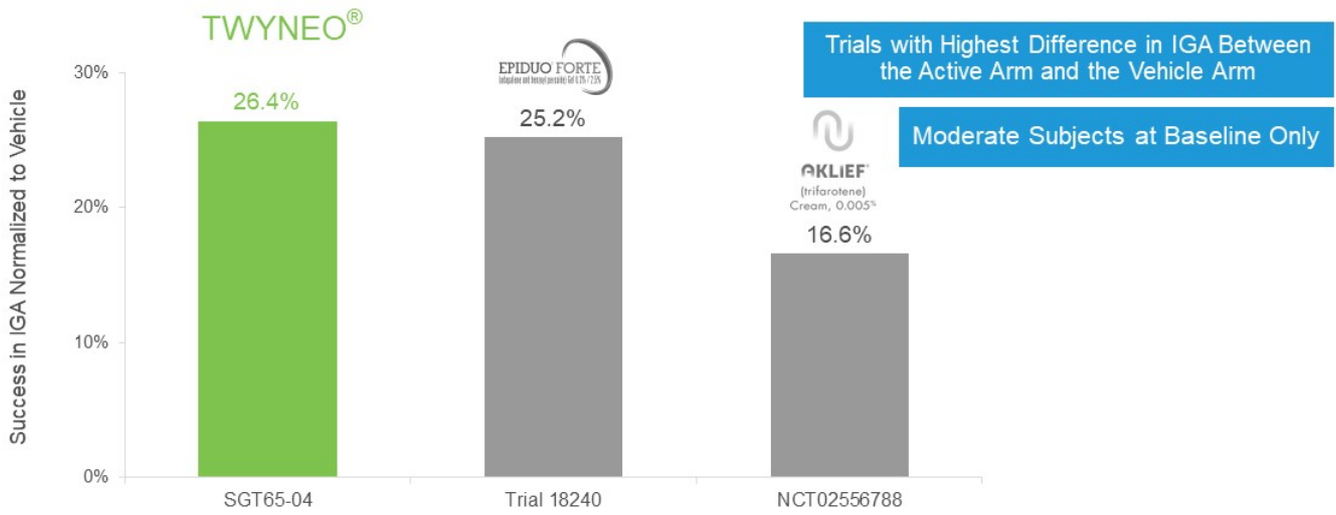
* Individual results vary



SUCCESS IN IGA



*Sol-Gel did not conduct a head-to-head comparison trial or study. The results described above are for illustrative purposes only and should not be construed as conclusions to be drawn as if we conducted a head-to-head comparison trial or study



*Sol-Gel did not conduct a head-to-head comparison trial or study. The results described above are for illustrative purposes only and should not be construed as conclusions to be drawn as if we conducted a head-to-head comparison trial or study



PRIMARILY MILD-TO-MODERATE

TREATMENT-EMERGENT ADVERSE EVENTS

Subjects with Treatment-Emergent Adverse Events (TEAEs)	Study 65-04		Study 65-05	
	TWYNEO (n=274)	Vehicle (n=139)	TWYNEO (n=281)	Vehicle (n=138)
Treatment-Related Mild & Moderate TEAEs	46 (16.8%) [^]	2 (1.4%) [^]	39 (13.8%) [^]	3 (2.2%)
Treatment-Related Severe TEAEs	4 (1.5%) [^]	0	1 (0.4%) [^]	0
Not-Related TEAEs	19 (6.9%)	13 (9.4%)	27 (9.6%)	15 (10.9%)
Missing Subjects	0	0	1 (0.4%)	0
Not-Related Serious TEAEs	0	0	1 (0.4%) [†]	1 (0.7%) [‡]

[^]Most frequently reported adverse events being application site pain, dryness, erythema and exfoliation

[^]Two subjects with application site pain, a third subject with application site pain and exfoliation, and fourth subject with application site pruritus

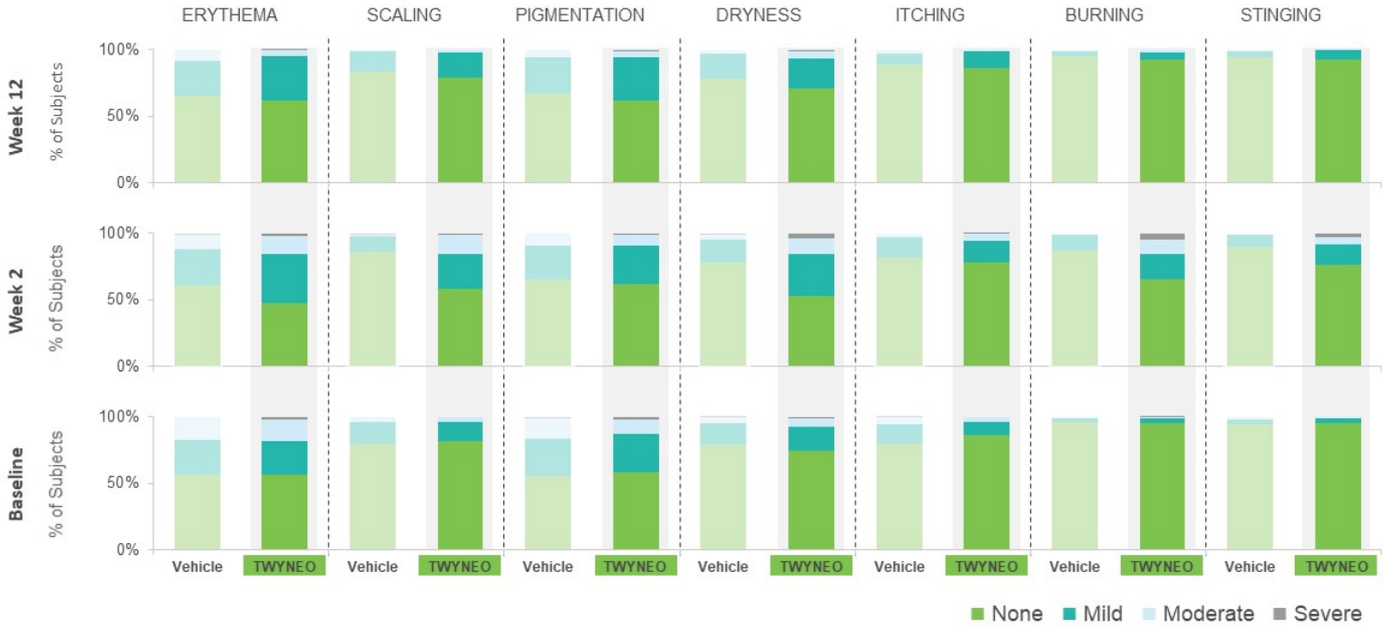
[^]One subject with application site pain, dryness and pruritus

[†]One subject with depression

[‡]One subject with depression, bipolar II disorder and conduct disorder



LOCAL SKIN REACTIONS





LOCAL SKIN REACTIONS





BROAD LONG-TERM
INTELLECUAL PROPERTY
ESTATE



- EPSOLAY is protected until 2032 by granted patents, and until 2040 by allowed patents
- TWYNEO is protected until 2038 by granted patents and until 2041 by pending patent applications
- 25 patent applications for erlotinib, tapinarof and roflumilast in various skin conditions (as of February 26, 2021)



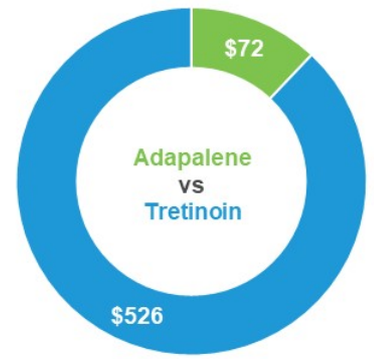
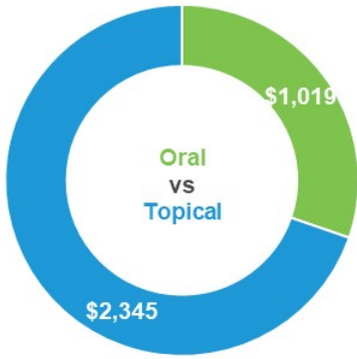
COMMERCIALIZATION
&
FINANCIALS

Branded Topicals are Important Segment



Source: IQVIA; Year 2019

**Branded Topical Combinations are Important Segment
 Tretinoin is the Most Prescribed Topical Retinoid**



Source: IQVIA; Year 2019



EPSOLAY & TWYNEO ARE COMPELLING ENOUGH TO DRIVE PAYOR COVERAGE

EPSOLAY[®]



- “All respondents recognized the product as a unique molecule for rosacea”
- “Near unanimous recognition as additional option for rosacea”
- “If priced and rebated similarly to the covered products, coverage seems likely”

TWYNEO[®]



- “Unique MOA will qualify it for formulary addition, price will determine its position”
- “If you price it like Epiduo, it will be managed like Epiduo”
- “If similarly priced with better tolerability, it would become preferred brand”

Sources: NaviSync LLC (Morristown, NJ), Sol-Gel Managed Market Access for Acne and Rosacea, July 2019
NaviSync LLC (Morristown, NJ), Twyneo Payer Market Research Topline Summary, February 2020



PRUDENT COMMERCIALIZATION APPROACH

We are in advanced negotiations with a potential partner regarding the commercialization of EPSOLAY® and TWYNEO®



* EPSOLAY & TWYNEO are investigational. Safety and efficacy have not been established

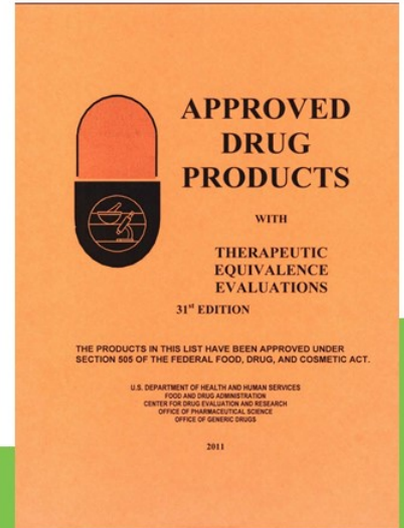
Source: Syneos Health (Morrisville, NC), Sol-Gel Market Analysis, June 2019



LUCRATIVE GENERIC PIPELINE

12 collaborations with Perrigo – with 50/50 gross profit sharing

- In June 2021, Perrigo launched ivermectin cream, 1%
- In February 2019, Perrigo launched acyclovir cream, 5%
- In June 2020, Perrigo was first-to-file a Paragraph IV Certification for Duobrii®
- In January 2020, Perrigo filed a Paragraph IV Certification for Bryhali®





FINANCIAL PROFILE

- Gross proceeds of \$86.3 million raised in IPO on February 5, 2018
- Gross proceeds of \$11.5 and \$23 million raised in public follow-on offerings on August 12, 2019, and February 13, 2020, respectively
- Additional \$5 million investment by controlling shareholder in April 2020
- 23,028,264 Ordinary Shares as of March 31, 2021
- \$8.7 million net revenues from generic products in 2020 and \$0.7 million net revenues from generic products in 1Q/21
- \$46.9 million in cash and investments as of March 31, 2021
- Under our operational model which assumes partnership regarding the commercialization of EPSOLAY® and TWYNEO® with a dermatology company that has a strong market presence, we expect that our cash resources will enable funding of operational and capital expenditure requirements into the third quarter of 2022



LOOKING FORWARD

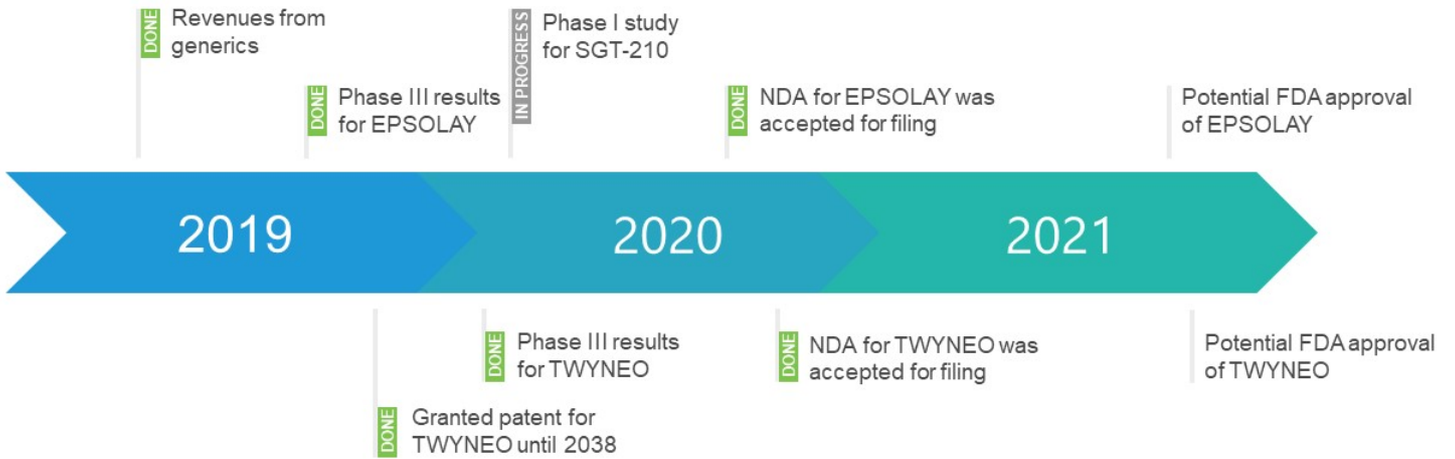
PALMOPLANTAR KERATODERMA



Palmoplantar keratoderma (PPK) is a group of skin conditions characterized by thickening of the skin on the palms of the hands and soles of the feet



RECENT MILESTONES & NEXT STEPS





Sol-Gel

Advanced Topical Therapy

NASDAQ: SLGL