UNITED STATES SECURITIES AND EXCHANGE COMMISSION

Washington, D.C. 20549

FORM 8-K

CURRENT REPORT

Pursuant to Section 13 or 15(d) of the Securities Exchange Act of 1934.

Date of Report: August 18, 2020 (Date of earliest event reported)

Oragenics, Inc.

(Exact name of registrant as specified in its charter)

FL (State or other jurisdiction of incorporation)

> 4902 Eisenhower Boulevard, Suite 125 Tampa, FL (Address of principal executive offices)

001-32188 (Commission File Number) 59-3410522 (IRS Employer Identification Number)

33634 (Zip Code)

813-286-7900 (Registrant's telephone number, including area code)

(Former Name or Former Address, if changed since last report)

Check the appropriate box below if the Form 8-K filing is intended to simultaneously satisfy the filing obligation of the registrant under any of the following provisions:

[] Written communications pursuant to Rule 425 under the Securities Act (17 CFR 230.425)

[] Soliciting material pursuant to Rule 14a-12 under the Exchange Act (17 CFR 240.14a-12)

[] Pre-commencement communications pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))

[] Pre-commencement communications pursuant to Rule 13e-4(c) under the Exchange Act (17 CFR 240.13e-4(c))

Securities registered pursuant to Section 12(b) of the Act:

Title of each class	Trading Symbol(s)	Name of each exchange on which registered
Common Stock	OGEN	NYSE American

Indicate by check mark whether the registrant is an emerging growth company as defined in Rule 405 of the Securities Act of 1933 (§230.405 of this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (§240.12b-2 of this chapter).

Emerging growth company []

If an emerging growth company, indicate by check mark if the registrant has elected not to use the extended transition period for complying with any new or revised financial accounting standards provided pursuant to Section 13(a) of the Exchange Act. []

Item 7.01 Regulation FD Disclosure.

Oragenics, Inc. ("Oragenics" or the "Company") will be making an investor presentation (the "Investor Presentation") on August 18, 2020. The Company also expects to use the Investor Presentation from time to time thereafter in connection with presentations to potential investors, industry analysts and others. A copy of the Investor Presentation is attached hereto as Exhibit 99.1 and is incorporated herein by reference. Additionally, the Investor Presentation will be available under the "Presentations" tab in the "News and Media" section of the Company's website, located at <u>www.oragenics.com</u>.

By filing this Current Report on Form 8-K and furnishing the information contained herein, the Company makes no admission as to the materiality of any information in this report that is required to be disclosed solely by reason of Regulation FD.

The information contained in the Investor Presentation is summary information that is intended to be considered in the context of the Company's Securities and Exchange Commission ("SEC") filings and other public announcements that the Company may make, by press release or otherwise, from time to time. The Company undertakes no duty or obligation to publicly update or revise the information contained in this report, although it may do so from time to time as its management believes is warranted. Any such updating may be made through the filing of other reports or documents with the SEC, through press releases or through other public disclosure.

The information presented in Item 7.01 of this Current Report on Form 8-K and Exhibit 99.1 shall not be deemed to be "filed" for purposes of Section 18 of the Securities Exchange Act of 1934, as amended (the "Exchange Act"), or otherwise subject to the liabilities of that section, unless the Company specifically states that the information is to be considered "filed" under the Exchange Act or specifically incorporates it by reference into a filing under the Securities Act of 1933, as amended, or the Exchange Act.

Item 9.01 Financial Statements and Exhibits.

(d) Exhibits.

Exhibit No.	Description
99.1	Investor Presentation.

SIGNATURES

In accordance with the requirements of the Exchange Act, the registrant caused this report to be signed on its behalf by the undersigned, thereunto duly authorized on this 18th day of August, 2020.

ORAGENICS, INC. (Registrant)

BY: /s/ Michael Sullivan Michael Sullivan

Chief Financial Officer



Developing Vaccines & Novel Antibiotics to Treat Tomorrow's Infections Corporate Presentation

NYSE American: OGEN

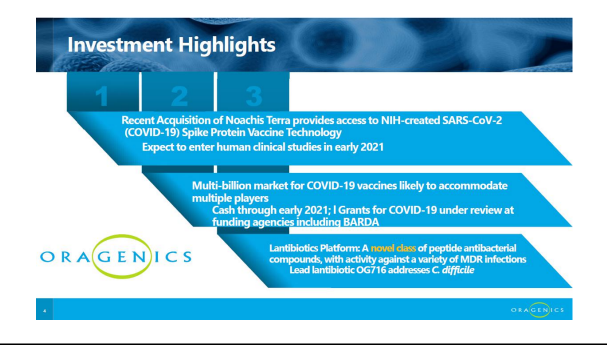
August 18, 2020

Safe Harbor Statement

Certain statements made in this presentation include forward-looking actions that Oragenics, Inc. ("Oragenics," or the "Company") anticipates based on certain assumptions. These statements are indicated by words such as "expect", "anticipate", "should" and similar words indicating uncertainty in facts, figures and outcomes. Such statements are made pursuant to the Safe Harbor Provisions of the Private Securities Litigation Reform Act of 1995. While Oragenics believes that the expectations reflected in such forward-looking statements are reasonable, it can give no assurance that such statements will prove to be correct. The risks associated with the Company are detailed in the Company's various reports filed by the Company with the Securities and Exchange Commission.

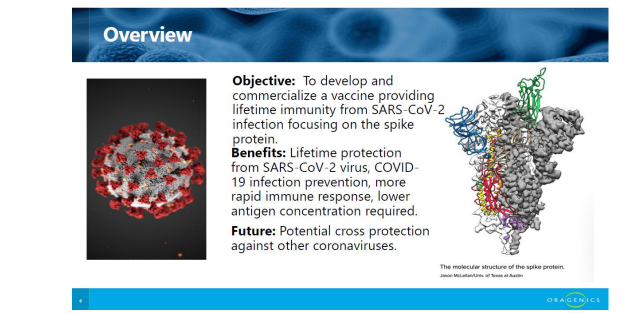
Oragenics Company Description

Oragenics (NYSE American: OGEN) is a development stage company dedicated to fighting infectious diseases. It is focused on advancing its TerraCoV2 vaccine candidate to combat the novel coronavirus pandemic, leveraging coronavirus spike protein research licensed from the National Institutes of Health. It is also developing lantibiotics, a novel class of antibiotic, focused on combatting multidrug-resistant organisms.





Lead Program: NIH-created SARS-CoV2 S-2P (COVID 19) vaccine candidate focused on the stabilized "Spike Protein"



Technology Overview – RSV F Protein: Spike Proteins Continually Changing Shape

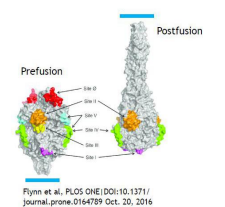
Accessible antigenic sites depend on protein conformation:

- Pre sites: O, V
- Post site¹: I
- Pre/post sites: II, III, IV

Absorption of human convalescent serum with postfusion F modestly reduces neutralizing antibodies

Absorption of human convalescent serum with prefusion F removes almost all neutralizing antibodies

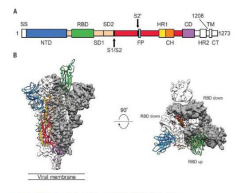
1. MAbs to site 1 preferentially bind postfusion conformation



Technology Overview – Enhanced Immune Response Utilizing Stabilized Spike Protein Antigen

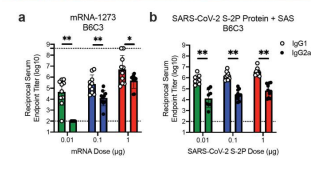
Stabilized Prefusion Spike Protein Ectodomain Trimer

- Class I fusion protein
- Two amino acid substitutions stabilize prefusion conformation
- T4 fibritin trimerization domain
- Expressed in mammalian cell line



Daniel Wrapp et al. Science 2020; 367:1260-1263

SARS-CoV-2 S-2P IgG subclass Results: B6C3 Mice



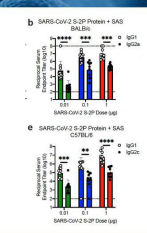
 Immunizations with mRNA-1273 and S-2P protein, delivered with TLR4 agonist, elicit S-specific Th1-biased T cell responses. B6C3F1/J mice were immunized at lgG1 weeks 0 and 3 with 0.01, 0.1, or 1 µg of mRNA-1273 lgG2a/c or SAS-adjuvanted SARS-CoV-2 S-2P protein. Sera were collected 2 weeks post-boost and assessed by ELISA for SARS-CoV-2 S-specific IgG1 and IgG2a/c.

Endpoint titers (a-b) were calculated. For mice for which endpoint titers did not reach the lower limit of detection (dotted line), ratios were not calculated (N/A).

SARS-CoV-2 S-2P protein with SAS adjuvant produces significant and balanced lgG1 and $lgG2_{a/c}$ levels at doses as low as 0.01 ug demonstrating level of immune response.

Corbett et al, BioRxiv. 2020. https://www.biorxiv.org/content/10.1101/2020.06.11.145920v1

SARS-CoV-2 S-2P IgG subclass Results: BALB/c and C57BL/6 Mice



SAS-adjuvanted S-2P protein elicit both IgG2a and IgG1 subclass S-binding antibodies. BALB/cJ (b) or C57BL/6J (e) mice were immunized at weeks 0 and 3 with 0.01 (green), 0.1 (blue), or 1 μ g (red) of SARSCOV- 2 S-2P protein adjuvanted with SAS. Sera were collected 2 weeks post-boost and assessed by ELISA for SARS-CoV-2 S-specific IgG1 and IgG2a or IgG2c for BALB/cJ and C57BL/6J mice, respectively.

Similar significant and balanced IgG1 and $IgG2_a$ or $IgG2_c$ titers were also observed in BALB/c and C57BL/6 mice immunized with SARS-CoV-2 S-2P protein with SAS adjuvant indicating the immune response is robust and seen in a second strain of mouse.

ORAGENICS

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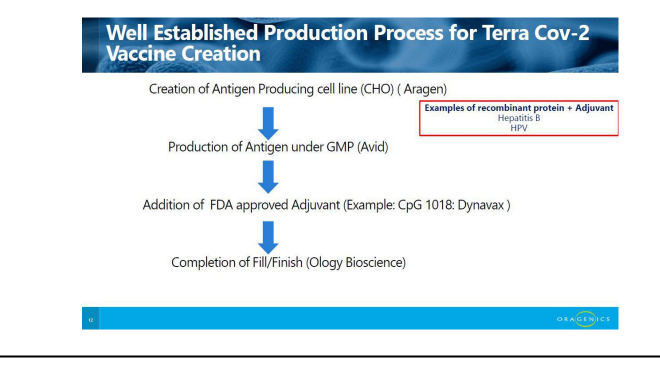
SARS-CoV-2 S-2P Neutralization Titers

Extended Data Table 1. Concordance of Pseudovirus Neutralization Assay and PRNT.

Mouse Serum	Reciprocal IC ₅₀ T	Fold	
Pool #1	Pseudovirus Neutralization ²	PRNT ³	Difference ⁴
1	893.5 +/- 1.4	933.5	1.0
2	211.6 +/- 1.5	314.5	0.7
3	159.8 +/- 1.3	397.1	0.5

NORMOV TINCE were immunized at weeks 0 and 3 with 1 µg SARS-CoV-2 S-2P protein, adjuvanted with SAS. Sera were collected 2
weeks post-boost and pooled (N = 3 mice/pool).
2. ICS0 titers were averaged from pseudovirus neutralization assays completed in 5 experimental replicates. (GMT +/- geometric SD)
3. ICS0 titer from PRNT assay completed once.
4. Fold difference calculated as average pseudovirus neutralization ICS0 titer relative to PRNT ICS0
Titer

SARS-CoV-2 S-2P protein with SAS adjuvant produces significant Nab titers in BALB/c mice.



Technology Overview – Current Status

- NIH/NIAID license secured with acquisition of Noachis Terra
- · Non-dilutive grants, including BARDA submitted or under development
- Utilize NIAID pre-clinical services for pre-IND enabling studies
- Contracts in place for:
 - Cell bank manufacture
 - Vaccine manufacture
 - Clinical Research Organization & Regulatory Consultants
- Utilize BARDA services for fill/finish
- Creation of cell line complete
- Development of analytical methods and transfer to manufacturing facility underway

Regulatory Strategy: Aggressive Fast Track Approach





Novel Lantibiotic Platform for Multidrug Resistant Bacterial Infections

2019 CDC List of Antibiotic Resistant Bacteria and Fungi

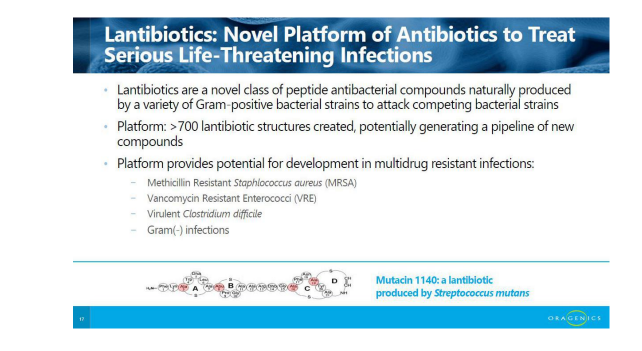
Urgent Threats

Carbapenem-resistant Acinetobacter
 Candida auris
 Clostridioides difficile
 Carbapenem-resistant Enterobacteriaceae
 Drug-resistant Neisseria gonorrhoeae
 Serious Threats
 Drug-resistant Campylobacter
 Drug-resistant Candida
 ESBL-producing Enterobacteriaceae
 Vancomycin-resistant Enterococci (VRE)
 Multidrug-resistant Pseudomonas aeruginosa
 Drug-resistant nontyphoidal Salmonella
 Drug-resistant Shigella
 Methicillin-resistant Staphylococcus
 aureus (MRSA)
 Drug-resistant Streptococcus pneumoniae
 Drug-resistant Streptococcus pneumoniae

Concerning Threats

Enythromycin-Resistant Group A Streptococcus
 Clindamycin-resistant Group B Streptococcus
 Watch List
 •Azole-resistant Aspergillus fumigatus

•Drug-resistant Mycoplasma genitalium •Drug-resistant Bordetella pertussis



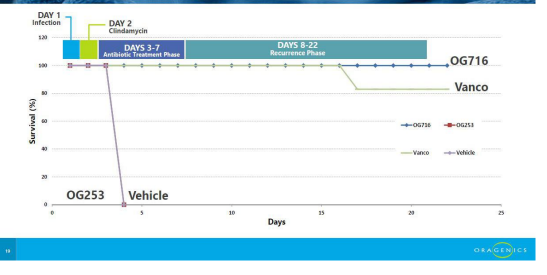
C. difficile and C. difficile Infection (CDI): Epidemiology

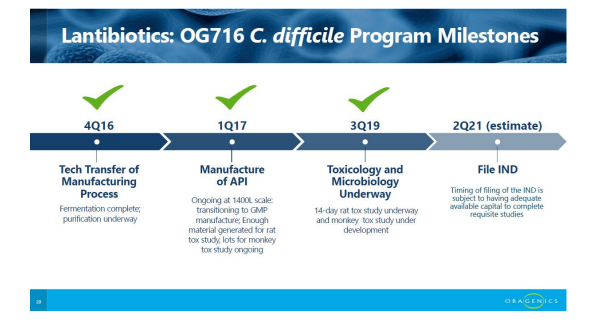
- *C. difficile* is an infection of the colon causing colitis by producing toxins that damage lining of the colon
- 223,900 infections annually resulting in 12,800 deaths
- 83,000 will experience at least one recurrence
- Healthcare-associated infections occur: 37% hospital onset, 36% nursing home onset, and 27% community onset
- C. *difficile* associated diarrhea is associated with a 1-2 week hospital stay at a cost of \$1BN/year
- **Emerging problem**: 8% of CDI associated with onset of concomitant Vancomycin Resistant Enterococci (VRE) infection





Oral OG716 Superior at Preventing C. difficile Deaths in Hamster Model









Capitalization

Common Stock Equivalents		
61,004,917	Cash	\$9.6M ⁽³⁾
2,261,703		
-		
20,513,145		
8,009,250		
91,789,015		
	Equivalents 61,004,917 2,261,703 - 20,513,145 8,009,250	Equivalents 61,004,917 Cash 2,261,703 - 20,513,145 8,009,250

(2) As of June 30, 2020, the Non-Voting, Non-Convertible Series C Preferred Shares have a stated value of \$33,847 per share and have an accruing of Preferred Shares resulted from the conversion of approximately \$3.3 million in debt obligations previously owed to Precigen (fka: Intrexon). The Series A, B, and C Preferred stock have no price based downround protection for the conversion price. 20% per ye

(3) Information is as of June 30, 2020 (excludes net proceeds from warrant exercises after the quarter).

Experienced Management Team

Dr. Alan F. Joslyn

Director, President and Chief Executive Officer

- Assumed CEO position at Oragenics in June 2016
- Held CEO positions at several private biotechnology companies including Sentinella Pharmaceuticals, Edusa Pharmaceuticals and Mt. Cook Pharma
- Over 25 years of drug development experience at Glaxo, Johnson & Johnson and Penwest

Mike Sullivan Chief Financial Officer

- Held senior-level financial positions for both publicly and privately held businesses
- Significant experience in product licensing and IP issues with strong background in both domestic and international retail operations

Dr. Martin Handfield

Senior Vice President, Discovery Research

- Molecular Microbiologist and former Tenured Associate Professor, College of Dentistry at The University of Florida
- Prolific researcher focusing on infectious diseases, host-pathogen interactions and non-invasive diagnostics

Experienced Management Team

Dr. David Zarley Consultant

- More than 30 years in vaccine research and development in the private sector
- Vice-President of Program Management for Vaccine Research and Development at Pfizer
- Senior Director /Medicines Team Leader for Pfizer Primary Care Business Unit
- Senior Director for Wyeth Research Project Management Business Unit
- Senior Director for Technical Operations and Product Supply (TOPS) at Wyeth Vaccines
- Senior Research Biochemist / Project Leader for Viral Vaccine Research and Development at Lederle-Praxis Biologicals

Near Term Milestones



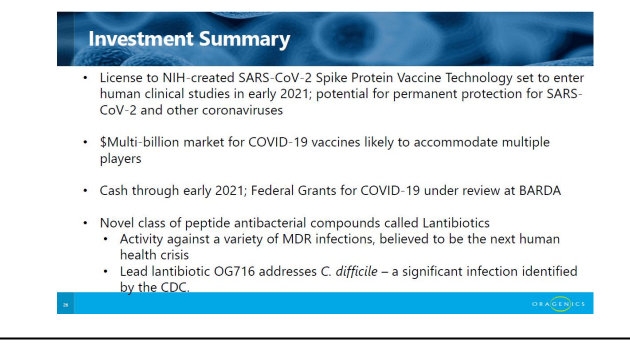
3Q20

- Complete CHO cell line Vaccine Development
- Initiate Vaccine CDMO work
- Complete FDA pre-IND meeting for Vaccine Candidate

4Q20

- Complete pre-clinical studies for Vaccine Candidate
- Advance Vaccine GMP manufacture at CDMO
- Initiate building of CHO MCB for the Vaccine
- 1Q21
- File IND for Vaccine
- 2Q21
- Initiate Phase 1 Clinical Study
- File IND for OG716





Oragenics, Inc. 4902 Eisenhower Blvd., Suite 125





NYSE American: OGEN

August 18, 2020