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 (Date of earliest event reported): May 14, 2021

VIRIOS THERAPEUTICS, INC.

(Exact Name of Registrant as Specified in Charter)

Delaware001-3981145-4618270(State or other jurisdiction
of incorporation)(Commission
File Number)(IRS Employer
Identification No.)

44 Milton Avenue Alpharetta, GA (Address of principal executive offices)

30009 (Zip Code)

Registrant's telephone number, including area code: (866) 620-8655

(Former name or former address, if changed since last report): Not Applicable

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, par value \$0.0001	mmon Stock, par value \$0.0001 VIRI Nasdaq Capita			

	the appropriate box below if the Form 8-K filing is intended to simultaneously satisfy the filing obligation of the registrant under any e 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))
(§23) If an	eate by check mark whether the registrant is an emerging growth company as defined in Rule 405 of the Securities Act of 1933 0.405 of this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (§240.12b-2 of this chapter).

Item 7.01 Regulation FD Disclosure.

On May 14, 2021, Virios Therapeutics, Inc. (the "Company") will be posting a presentation to its website that may be used by the Company from time to time with investors, analysts, collaborators, vendors or other third parties. A copy of the presentation is furnished as Exhibit 99.1

The information in this Item 7.01, including the attached exhibit, is furnished solely pursuant to Item 7.01 of Form 8-K. Consequently, such information is not deemed "filed" for purposes of Section 18 of the Securities Exchange Act of 1934, or otherwise subject to the liabilities of that section. Further, the information in this Item 7.01, including the exhibit, shall not be deemed to be incorporated by reference into the filings of the registrant under the Securities Act of 1933.

Cautionary Statement Regarding Forward-Looking Information

This current report on Form 8-K contains "forward-looking statements" within the meaning of the Private Securities Litigation Reform Act of 1995. All statements other than those of historical fact in this presentation and accompanying oral commentary are forward-looking statements. Forward-looking statements may be identified by terminology such as "believe," "anticipate," "plan," "may," "intend," "will," "should," "expect," "estimate," "potential" and "continue" and similar expressions, including the negative of these words, but not all forward-looking statements contain these words. Forward-looking statements include, but are not limited to, statements regarding the Company's expectations regarding our future financial or business performance, plans, prospects, trends or strategies, objectives of management, competition and other financial and business matters; the potential, safety, efficacy, and regulatory and clinical progress of our current and prospective product candidates, planned clinical trials and preclinical activities, and projected research and development costs; the estimated size of the market for our product candidates; and the timing and success of our development and commercialization of our anticipated product candidates and the market acceptance thereof. Forward-looking statements are based on our current expectations and are subject to inherent uncertainties, risks and assumptions that are difficult to predict. Further, certain forward-looking statements are based on assumptions as to future events that may not prove to be accurate. 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 statements, including, but not limited to, the following: the outbreak of the novel coronavirus disease, COVID-19, has adversely impacted and may continue to adversely impact our business, including our preclinical studies and clinical trials; our limited operating history, which may make it difficult to evaluate our current business and predict our future success and viability; we have and expect to continue to incur significant losses; our need for additional funding, which may not be available; our substantial dependence on the success of our lead product candidate; failure to identify additional product candidates and develop or commercialize marketable products; the early stage of our development efforts; potential unforeseen events during clinical trials could cause delays or other adverse consequences; risks relating to the regulatory approval process or ongoing regulatory obligations; our product candidates may cause serious adverse side effects; our reliance on third parties; effects of significant competition; the possibility of system failures or security breaches; risks relating to intellectual property; our ability to attract, retain and motivate qualified personnel; and significant costs as a result of operating as a public company. These and other risks and uncertainties are described more fully in the section titled "Risk Factors" in the Annual Report on Form 10-K for the year ended December 31, 2020 filed with the Securities and Exchange Commission ("SEC") and elsewhere in our filings and reports with the SEC. These risks, uncertainties and other factors may cause our actual results to differ materially and adversely from what is contained in (or may be implied from) any forward-looking statements. Forward-looking statements speak as of the date they are made, and the Company undertakes no obligation to update them except as may be required under applicable law.

Item 9.01 Financial Statements and Exhibits.

<u>(d)</u>	Exhibits.		
Exhibit	Number	Description	
99.1		Presentation dated May 2021 (furnished herewith).	

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 hereunto duly authorized.

VIRIOS THERAPEUTICS, INC.

By: /s/ Angela Walsh

Name: Angela Walsh

Title: SVP of Finance and Corporate Secretary

May 14, 2021

Virios Therapeutics, Inc. (Nasdaq: "VIRI")

New Scientific Paradigm Exploring Herpes Simplex 1 (HSV-1) Virus Activation as Potential Underlying Cause of Fibromyalgia and Other Chronic Conditions

Corporate Update Q2 2021



HSV-1 virus

Forward Looking Statements

- Statements in this presentation contain "forward-looking statements" that are subject to substantial risks and uncertainties. Forward-looking statements contained in this presentation may be identified by the use of words such as "anticipate," "expect," "believe," "will," "may," "should," "estimate," "project," "outlook," "forecast" or other similar words, and include, without limitation, all statements other than those regarding historical facts, statements regarding Virios Therapeutics, Inc.'s expectations regarding our future financial or business performance, plans, prospects, trends or strategies, objectives of management, competition and other financial and business matters; the potential, safety, efficacy, and regulatory and clinical progress of our current and prospective product candidates, planned clinical trials and preclinical activities, and projected research and development costs; the estimated size of the market for our product candidates; and the timing and success of our development and commercialization of our anticipated product candidates and the market acceptance thereof. Forward-looking statements are based on our current expectations and are subject to inherent uncertainties, risks and assumptions that are difficult to predict. Further, certain forward-looking statements are based on assumptions as to future events that may not prove to be accurate. 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 statements, including, but not limited to, the following: the outbreak of the novel coronavirus disease, COVID-19, has adversely impacted and may continue to adversely impact our business, including our preclinical studies and clinical trials; our limited operating history, which may make it difficult to evaluate our current business and predict our future success and viability; we have and expect to continue to incur significant losses; our need for additional funding, which may not be available; our substantial dependence on the success of our lead product candidate; failure to identify additional product candidates and develop or commercialize marketable products; the early stage of our development efforts; potential unforeseen events during clinical trials could cause delays or other adverse consequences; risks relating to the regulatory approval process or ongoing regulatory obligations; our product candidates may cause serious adverse side effects; our reliance on third parties; effects of significant competition; the possibility of system failures or security breaches; risks relating to intellectual property; our ability to attract, retain and motivate qualified personnel; and significant costs as a result of operating as a public company. These and other risks and uncertainties are described more fully in the section titled "Risk Factors" in the Annual Report on Form 10-K for the year ended December 31, 2020 filed with the Securities and Exchange Commission ("SEC") and elsewhere in our filings and reports with the SEC. While we may elect to update these forward-looking statements at some point in the future, we assume no obligation to update or revise any forward-looking statements except to the extent required by applicable law. Although we believe the expectations reflected in such forward-looking statements are reasonable, we can give no assurance that such expectations will prove to be correct. Accordingly, readers are cautioned not to place undue reliance on these forward-looking statements. No representations or warranties (expressed implied) are made about the accuracy of any such forward-looking statements.
- This presentation also contains estimates and other statistical data made by independent parties and by us relating to market size and other data about our industry. This data involves a number of assumptions and limitations, and you are cautioned not to give undue weight to such data and estimates. In addition, projections, assumptions and estimates of our future performance and the future performance of the markets in which we operate are necessarily subject to a high degree of uncertainty and risk. Neither we nor our affiliates, advisors or representatives makes any representation as to the accuracy or completeness of that data or undertake to update such data after the date of this presentation.
- You should read the documents that we have filed with the SEC for more complete information about us. We encourage you to read such documents in full for more detailed information on statistics, reports and clinical trials referenced in this presentation. You may access these documents for free by visiting EDGAR on the SEC website at http://www.sec.gov.

Virios Therapeutics Summary



The Fibromyalgia (FM) Market is Large, but Dissatisfied – Highlighting Significant Commercial Potential for a Differentiated New FM Treatment



Our Lead Asset, Oral IMC-1, Demonstrated Significant FM Pain Reduction and Tolerability Benefits in Phase 2a Clinical Trial



Unique IMC-1 Antiviral Combination (famciclovir & celecoxib) Granted First Ever FDA Fastrack Review Designation for FM Treatment



Positive IMC-1 P2b Results in Q2 '22 Could Transform FM Patient Care



Virios Team and Board of Directors Have Led Development or Commercialization for Two of Three Approved FM Medicines



3

Proven Leadership Team with Experience in Fibromyalgia (FM) Development and Commercialization









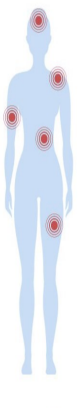
Fibromyalgia Disease Overview

Disease Characteristics

- FM is a Chronic Disease that Affects up to 8% of the US Population
- · Hallmark Characteristics are Widespread Chronic Pain and Severe Fatigue
 - Symptoms Present for ≥ 3 Months
- Other Symptoms May Include GI, Sleep, Mood Disorder and Headache

Devastating Impact

- Patients with FM > 3x Risk of Committing Suicide v. General Population
- · High Healthcare Utilization and Significant Disability
- Estimates Suggest as Many as 40% of FM Patients are Treated with Opioids
 - Opioid-treated Patients have Worse Outcomes Across Multiple Assessment Domains



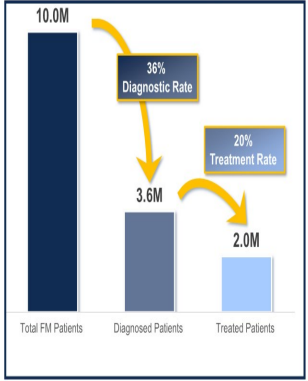


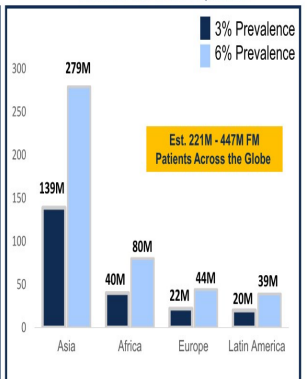


The Fibromyalgia Market is Large and Poised for Growth if Better Therapeutic Options Emerge

Significant US FM Market
Growth Potential Still Exists





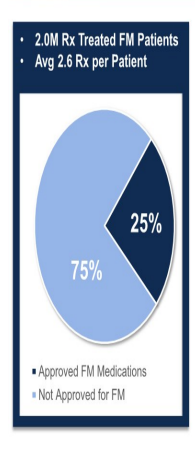


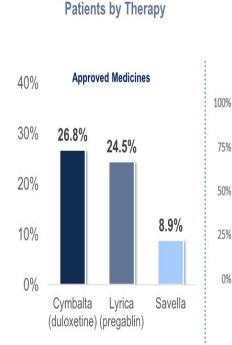
Source: National Fibromyalgia and Chronic Pain Association 2021; Vincent, A et al Arthritis Care Research 2013; Robinson et al Pain Medicine, 2012



Polypharmacy and Utilization of Unapproved Therapies Demonstrate Significant Unmet Need

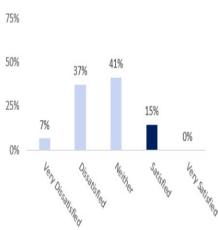
Share of US Treated FM





Only 15% of FM Treatment
Prescribers Reported Satisfaction
with Existing Treatments







7 Source: Robinson et al, *Pain Medicine*, 2012, . Lumleian primary research for Virios Therapeutics, Inc.

A New Treatment for Fibromyalgia Could Have Significant Commercial Potential in the US Market Alone

Total Projected US FM Community

FM Costs US \$12-14B Annually, 1-2% Lost Productivity in US

25% Patient Share of Treated FM Population @ 500K Unique Patients

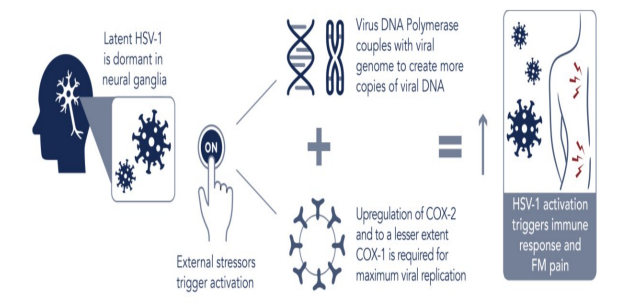
Lyrica @ \$16/day in 2019, Est. 60% Compliance Rate, Annual FM Sales Est. @ \$1.75B





Source: National Fibromyalgia and Chronic Pain Association, 2021; Evaluate Pharma, 2019; Robinson et al Pain Medicine 2012, Vincent et al Arthritis Care Research, 2013; Redbook, 2019

Discovery Implicates Dormant HSV-1 Reactivation Triggers Immune Response and Manifestation of Fibromyalgia

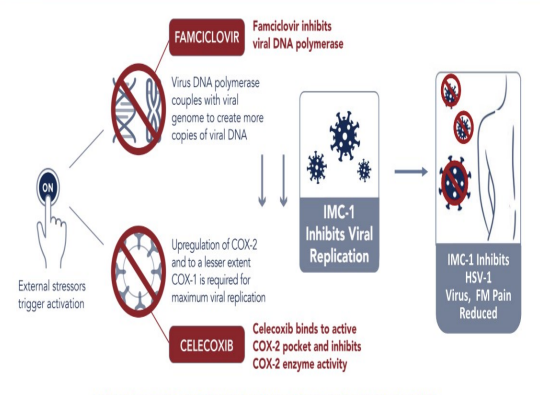


The World Health Organization Estimates That 3.7 Billion People Under the Age of 50 – or 67% of the Population are infected with Herpes Simplex Virus Type 1 (HSV-1), a Life-long Infection

Source: P.A. Bond, Medical Hypotheses, 1993,; R. A Vere Hodge and Y.-G. Cheng, Antiviral Chemistry & Chemotherapy, 1993; Kaufman et al, IOVS, 2005; Liu Y, et al, Scientific World Journal, 2014; Higaki S, et al Current Eye Research, 2009; Francisco Javier Ibañez et al, Frontiers in Microbiology, 2018,



IMC-1's Synergistic Antiviral Mechanism Serves as Basis for Proposed Fibromyalgia Treatment Effect



IMC-1 Composition of Matter Intellectual Property Protection to 2033

Source: P.A. Bond, Medical Hypotheses, 1993,; R. A Vere Hodge and Y.-G. Cheng, Antiviral Chemistry & Chemotherapy, 1993; Liu Y, et al, Scientific World Journal, 2014; Higaki S, et al Current Eye Research, 2009; Francisco Javier Ibañez et al, Frontiers in Microbiology, 2018



IMC-1: Completed Phase 2a Clinical Proof of Concept Trial

Design Summary:

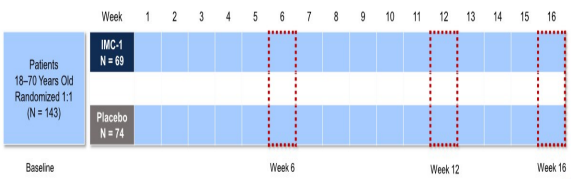
- · Randomized, Double-blind, Multi-center, Placebo-controlled
- IMC-1 (famciclovir + celecoxib) vs Placebo, Dosed Twice Daily
- Diagnosis of Fibromyalgia Using 2010 American College of Rheumatology, Assessments at Weeks 6, 12 and 16
- · 7-day Washout of FM Drugs and Opioids



Key Secondary Endpoints

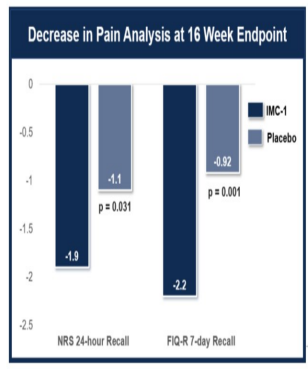
PGIC, FIQ-R Domains, 30% & 50% pain responder analyses

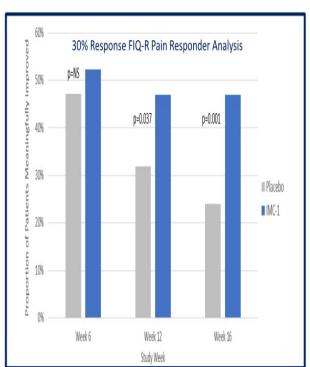
Received study drug treatment for a total of 16 weeks





IMC-1 Demonstrated Statistically Significant Reduction in Pain in Phase 2a Clinical Trial



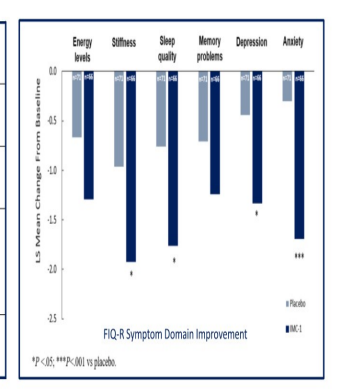


Source: Pridgen et al, Journal of Pain Research, 2017, Virios Therapeutics, Inc.



Key Secondary Outcomes at 16 Weeks

Endpoints	P Value	
PROMIS (NIH) Fatigue Assessment	p=0.001	
Patient's Global Impression of Change	P=0.040	
Fibromyalgia Impact Questionnaire	p=0.002	
Use of Rescue Medication	p=0.037	



Source: Pridgen et al, Journal of Pain Research, 2017, 2017, Virios Therapeutics, Inc.



IMC-1 Had a Lower Discontinuation Rate Versus Placebo in Fibromyalgia Phase 2a Study

	Placebo	IMC-1	IMC-1 Difference	
Discontinuation reasons:				
Adverse event (p=0.012)	12 (16.2%)	4 (5.8%)	2.8X reduction	
Therapeutic failure	12 (16.2%)	5 (7.2%)	2.3X reduction	
Other	5 (6.8%)	3 (4.4%)	1.5X reduction	



IMC-1 Phase 2b Design to be Conducted Using Optimized IMC-1 Dosage

Primary Endpoint: Reduction in pain

Secondary Endpoints: Fatigue, sleep, global health status, and patient functionality

FM Patients 2016 ACR criteria **Age** 18 - 65

Sample Size 460 (~230/arm) Treatments IMC-1 vs Placebo

Daily Assessments 16 Weeks

	2021			2022		
	Q1	Q2	Q3	Q4	Q1	Q2
Enrollment						
Study Duration						
P2b Topline Results						



IMC-1 Pipeline Potential Extends to Other Functional Somatic Syndromes

FIBROMYALGIA

- IMC-1 Statistically Significant Reduction in Pain P2a
- FDA Fast Track Review Designation, Progressing to P2b

IRRITABLE BOWEL SYNDROME

- Univ. of AL GI Biopsy Data Confirm Active HSV-1 in IBS
- ROME IV IBS Assessment Criteria Places Increasing Focus on Pain

CHRONIC FATIGUE SYNDROME/ MYALGIC ENCEPHALITIS

- Viral infections identified as possible triggers of CFS/ME
- Fatigue most common symptom beyond 2 weeks after COVID-19 onset
- IMC-1 Statistically Significant Reduction in fatigue in FM P2a



Capital Table as of May 12th 2021

VIRI S Therapeutics			
Listing/Symbol	Nasdaq/VIRI		
Common Shares Outstanding	8,330,390		
Underwriters Warrants* (Exercisable at \$12.50/share)	172,500		
Stock Options (Employees, Directors & Officers)**	874,397		
Fully Diluted Shares Outstanding	9,377,287		
Management Ownership on a Fully Diluted Basis	18.4%		

^{*}Become Exercisable on December 16, 2021



^{**}Weighted Avg. Exercise Price of \$9.80/share

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