

**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): January 8, 2024

**Cocrystal Pharma, Inc.**

(Exact name of registrant as specified in its charter)

Delaware  
(State or other jurisdiction  
of incorporation)

001-38418  
(Commission  
File Number)

35-2528215  
(IRS Employer  
Identification No.)

19805 N. Creek Parkway  
Bothell, WA  
(Address of principal executive offices)

98011  
(Zip Code)

Registrant's telephone number, including area code: (877) 262-7123

N/A  
(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 (see General Instruction A.2. below):

- ☐ 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	COCP	The Nasdaq Stock Market LLC (The Nasdaq Capital Market)

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.**

On January 8, 2024, Cocrystal Pharma, Inc. (the "Company") is making a presentation to certain members of the investment community. A copy of the slide presentation is being furnished as Exhibit 99.1.

The information in this Item 7.01 (including Exhibit 99.1) shall not be deemed "filed" for purposes of Section 18 of the Securities Exchange Act of 1934 (the "Exchange Act") or otherwise subject to the liabilities under such section, and shall not be deemed to be incorporated by reference into any filing of the Company under the Securities Act of 1933 or the Exchange Act.

**Item 9.01 Financial Statements and Exhibits.**

Exhibit No.	Description
99.1	<u>Cocrystal Pharma, Inc. Corporate Presentation, dated January 8, 2024</u>
104	Cover Page Interactive Data File (embedded within the Inline XBRL document)

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.

Dated: January 8, 2024

Cocrystal Pharma, Inc.

By: /s/ James Martin  
Name: James Martin  
Title: Chief Financial Officer and Co-Chief Executive Officer

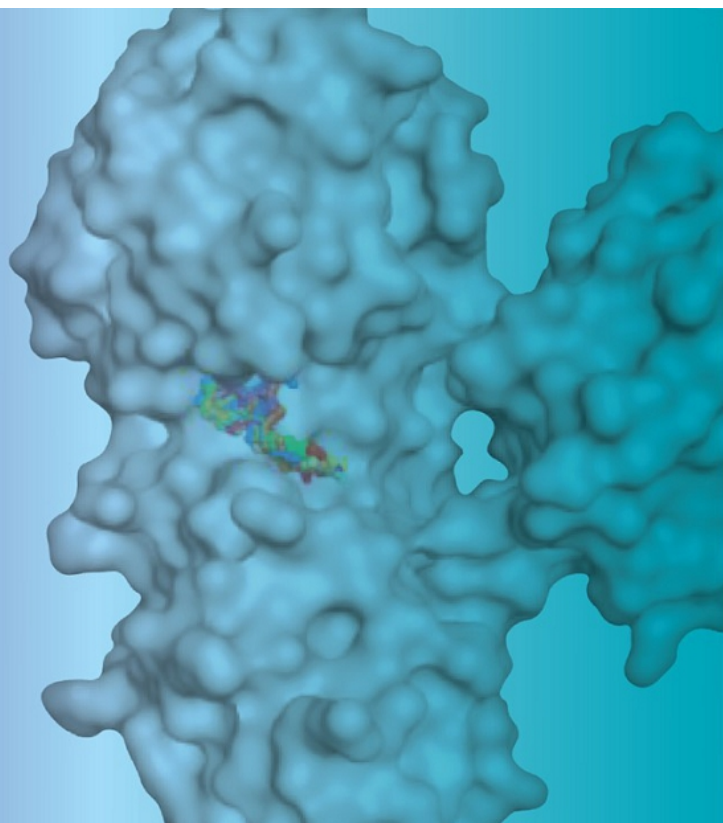
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Potent antivirals to combat some  
of the most serious diseases  
facing humanity

January 2024

Nasdaq: COCP  
[www.cocrystalpharma.com](http://www.cocrystalpharma.com)



## Forward-Looking Statements

This presentation contains forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995, including statements regarding the market opportunities for the treatment of acute and chronic viral diseases which are the focus of our programs; our development pipeline; our technology platform's ability to produce viable drug candidates at reduced development timelines and costs; the expected future characteristics and progress of product candidates and development efforts in our clinical programs, including our ongoing Phase 2a study for oral influenza PB2 inhibitor; our ongoing Phase 1 study with 3CL protease inhibitor for coronavirus and norovirus; and a planned Phase 1 study for inhaled influenza PB2 inhibitor this year; our exploration of other collaboration opportunities including our pursuit of opportunities related to pandemic preparedness, and the expected sufficiency of our cash balance to fund our planned operations.

Forward-looking statements are prefaced by words such as "anticipate," "expect," "plan," "could," "may," "will," "should," "would," "intend," "seem," "potential," "appear," "continue," "future," "believe," "estimate," "forecast," "project," and similar words. Forward-looking statements are based on our current expectations and assumptions regarding our business, the economy and other future conditions. Because forward-looking statements relate to the future, they are subject to inherent uncertainties, risks and changes in circumstances that are difficult to predict. We caution you, therefore, against relying on any of these forward-looking statements. Our actual results may differ materially from those contemplated by the forward-looking statements for a variety of reasons, including, without limitation, the risks arising from interest rate increases in response to inflation, uncertainty in the financial markets, the possibility of a recession and the geopolitical conflict in Israel and Ukraine on our Company, our collaboration partners, and on the U.S., U.K., Australia and global economies, our ability to proceed with studies including recruiting volunteers for and procuring or manufacturing materials for such studies by our clinical research organizations and vendors, the results of the Phase 2a and Phase 1 studies referred to above, our and our collaboration partners' technology and software performing as expected and maintenance and protection of related intellectual property rights, financial difficulties experienced by certain partners and our ability to secure and maintain new collaboration partners, the results of any current and future preclinical and clinical trials, general risks arising from clinical trials, receipt of regulatory approvals, regulatory changes, development of effective treatments and/or vaccines by competitors, including as part of the programs financed by the U.S. government, and potential mutations in the viruses we are targeting which may result in variants that are resistant to a product candidate we develop. Further information on our risk factors is contained in our filings with the Securities and Exchange Commission, including our Annual Report on Form 10-K for the year ended December 31, 2022. Any forward-looking statement made by us in this presentation speaks only as of the date on which it is made. Factors or events that could cause our actual results to differ may emerge from time to time, and it is not possible for us to predict all of them. We undertake no obligation to publicly update any forward-looking statement, whether as a result of new information, future developments or otherwise, except as may be required by law.

*Applying powerful, proprietary drug discovery platform technology to develop first- and best-in-class broad-spectrum antiviral drugs*

### Advancing programs in high-value antiviral drug targets

- Pandemic and seasonal influenza
- Pandemic norovirus
- Pandemic coronavirus and respiratory viruses

### Drug candidates with clinically validated mechanisms of action

- Effectively cure viral diseases
- Broad-spectrum and potent antiviral activity
- Designed to be effective for emerging variants and existing drug resistant viruses
- Multiple routes of administration (oral, inhalation, and injectable)

### Proprietary drug discovery platform technology

- Unique drug discovery platform technology developed with Nobel Prize-winning technology

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## Investment Highlights

- Targeting multibillion-dollar, global markets for the treatment of acute and pandemic viral diseases
- Proprietary structure-based drug-discovery platform technology provides opportunity for discovery and development of novel, potent, broad-spectrum drug candidates
- Advancing multiple clinical programs
  - Oral influenza PB2 inhibitor CC-42344 – Phase 2a initiated in 2023
  - First dual oral coronavirus-norovirus protease inhibitor CDI-988 – Phase 1 initiated in 2023
  - Inhaled influenza PB2 inhibitor CC-42344 – Phase 1 to begin in 2024
- Developing multiple discovery programs for respiratory viral diseases
  - Pan-viral protease inhibitors
  - Influenza replication inhibitors
- Additional pandemic preparedness collaboration opportunities are being explored
- Seasoned leadership includes experienced management, senior scientists and two Nobel laureates
- Cost-efficient operations and clean capital structure; cash sufficient to fund planned operations

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## Multiple clinical assets poised to deliver significant growth

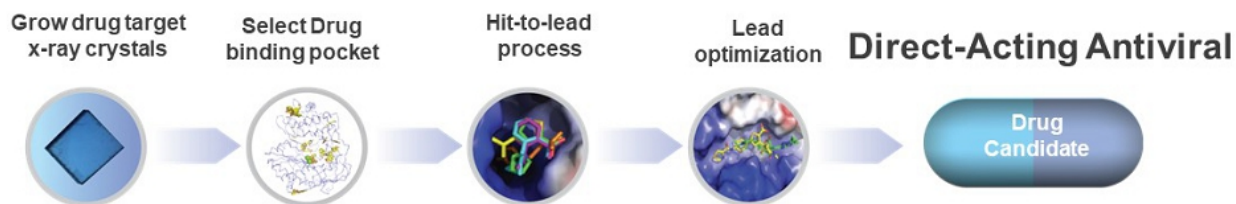
Program	Candidate	Discovery	Preclinical	Phase 1	Phase 2	Phase 3
Influenza A	Oral PB2 inhibitor CDI-988					Phase 2a initiated in 2023
	Inhaled PB2 inhibitor CDI-988					Phase 1 study planned in 2024
Influenza A/B	Replication inhibitors					Lead optimization ongoing
Norovirus & Coronavirus	Oral Pan-viral protease inhibitor CDI-988					Phase 1 initiated in 2023
Coronavirus (Licensed)	Protease inhibitor CDI-45205					IND-enabling study ongoing
Norovirus	Replication inhibitors					Discovery ongoing
Respiratory viruses	Pan-viral inhibitors					Discovery ongoing

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## Proprietary Drug Discovery Platform Technology for Direct-Acting Antivirals

Cocrystal's technology platform provides potential for novel drug candidates at reduced development timelines and costs

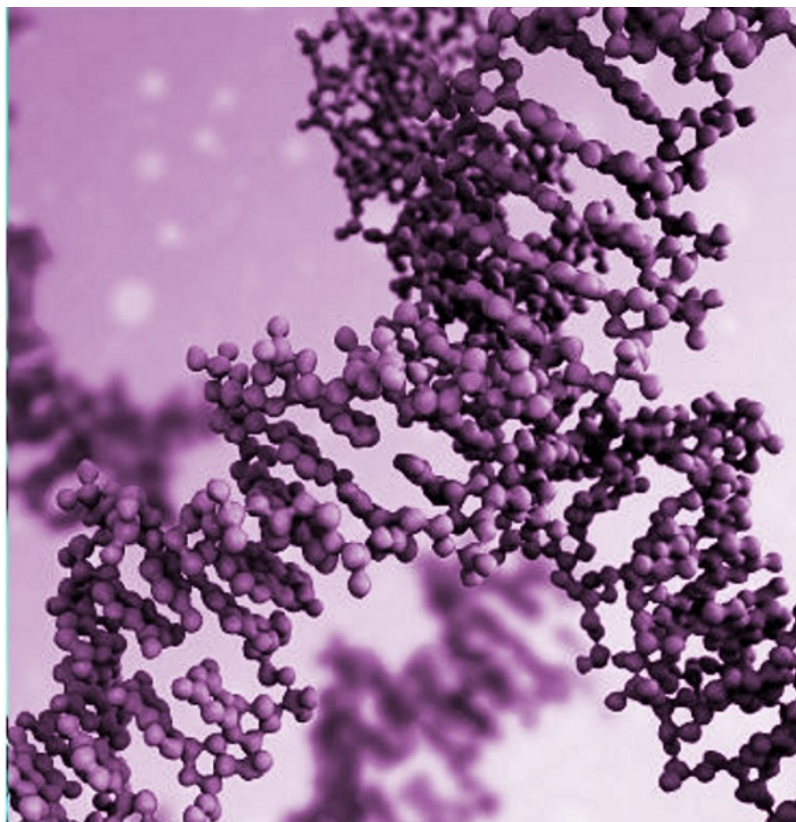


Provide high-resolution 3D structures of drug target complexed with inhibitor at atomic level

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# Pandemic and Seasonal Influenza Program



## Pandemic and Seasonal Influenza: A Major Global Health Concern

- 1 billion cases, 3-5 million severe illnesses and up to 650,000 deaths worldwide annually<sup>1</sup>
- Not well managed with currently approved vaccines having only 40-60% effectiveness<sup>2</sup>
- On average ~8% of the U.S. population contracts influenza each season<sup>3</sup>
- Influenza is responsible for ~\$10.4 billion in direct costs for hospitalizations and outpatient visits for adults in the U.S. annually
- Only influenza A causes pandemic flu and is also responsible for the majority of seasonal influenza infections<sup>1</sup>
- Potential emerging pandemic influenza A strains and drug resistant strains against approved influenza antivirals, Tamiflu® and Xofluza®
  - Tamiflu® has long history of drug resistance<sup>5</sup>
  - Xofluza® has shown emergence of drug resistant mutations<sup>6</sup>

<sup>1</sup> World Health Organization (WHO) (March 2019): [https://www.who.int/news-room/fact-sheets/detail/influenza-\(seasonal\)](https://www.who.int/news-room/fact-sheets/detail/influenza-(seasonal))

<sup>2</sup> Center for Disease Control and Prevention (CDC): Vaccine Effectiveness: How Well Do Flu Vaccines Work?: <https://www.cdc.gov/flu/vaccines-work/vaccineeffect.htm>

<sup>3</sup> CDC Seasonal Flu Microsite

<sup>4</sup> CDC: Make It Your Business to Fight the Flu

<sup>5</sup> ScienceDaily (March 2014) Tamiflu-resistant influenza related to mutations in genome: <https://www.sciencedaily.com/releases/2014/03/140331114237.htm>

<sup>6</sup> NEJM Journal Watch (September 2018) A Promising Drug for Influenza?: <https://www.jwatch.org/na47413/2018/09/12/promising-drug-influenza>

# Influenza Development Programs Focused on Therapeutic and Prophylactic Replication Inhibitors

## Clinical assets for pandemic and seasonal influenza

### Oral PB2 inhibitor CC-42344

- Ongoing Phase 2a
- Potent broad-spectrum activity
- Favorable safety profile and tolerability
- Potential for best-in-class

### Inhaled PB2 inhibitor CC-42344

- Ongoing GLP tox study
- Potent broad-spectrum activity
- Superior pulmonary exposure
- Potential for both prophylactic and therapeutic treatments

## Promising Early-Stage Programs

### Replication inhibitors

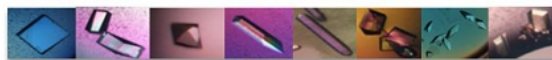
- Discovery ongoing
- Potent broad-spectrum activity against influenza A and B strains
- Novel mechanisms of action

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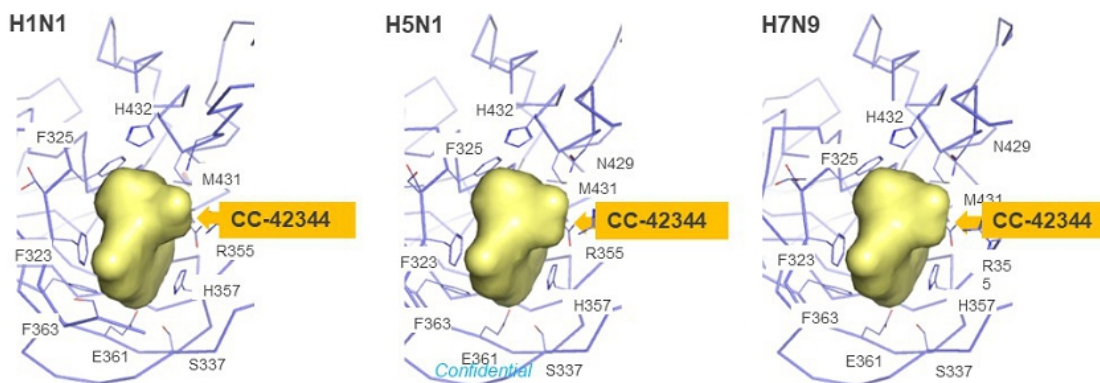
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## CC-42344 Binds to Highly Conserved Active Site of Influenza A PB2 Protein

### Cocrystal proprietary drug discovery platform technology

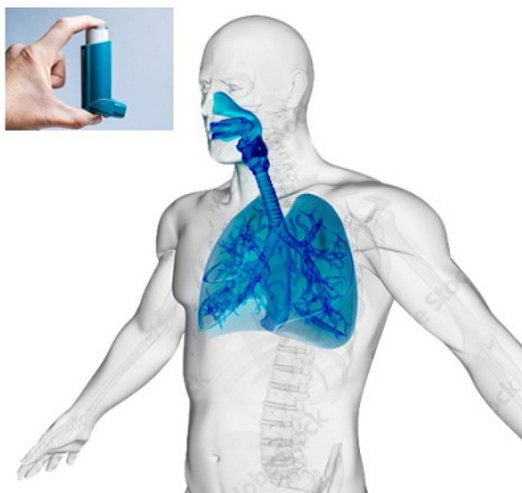


### Highly pathogenic influenza A strains



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## Advantages of inhalation antiviral therapy

- Directly targets infected respiratory epithelial cells
- Achieves higher accumulation of drug in the pulmonary system
- Produces rapid clinical response
- Reduces potential systemic side effects

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## CC-42344 Shows Potent Antiviral Activity in Influenza-Infected Lung Epithelium

### Uninfected human bronchial airway epithelia



Influenza A  
H1N1 infection

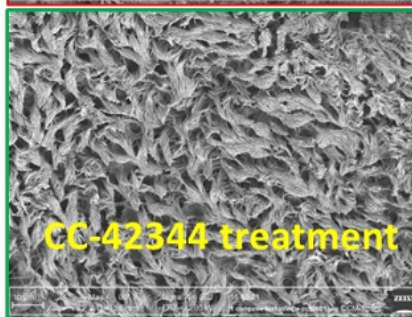


Cells are killed by H1N1 virus, and most cilia are destroyed

No CC-42344



CC-42344 treatment



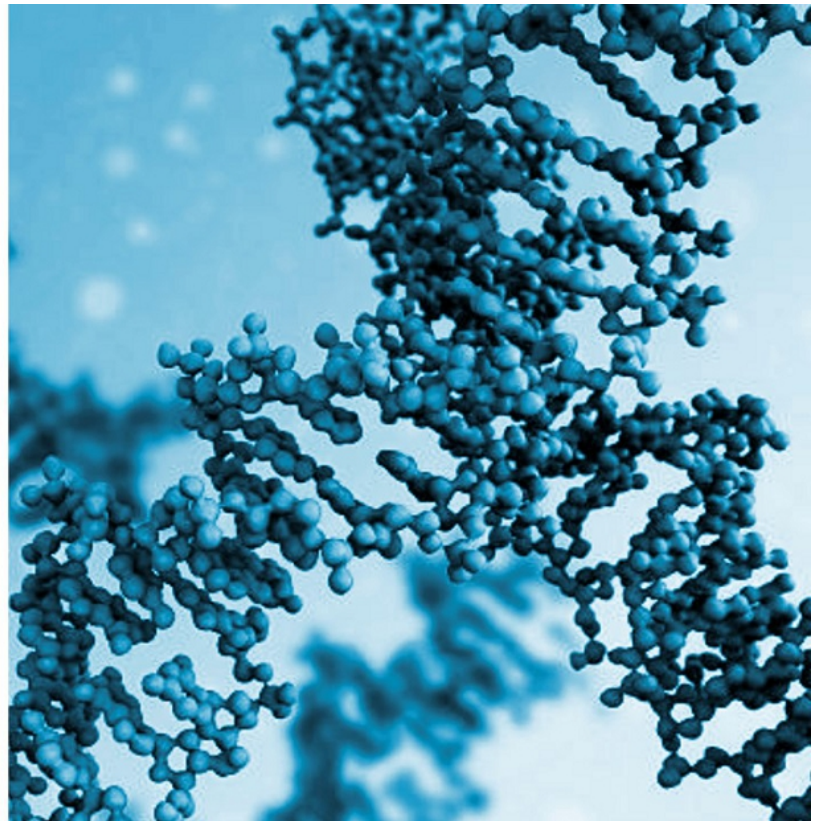
- Favorable safety profile: No toxicity in CC-42344-treated human lung epithelium
- Showed potent antiviral activity in influenza A (H1N1)-infected human lung epithelium
- Inhalation formulation development is completed

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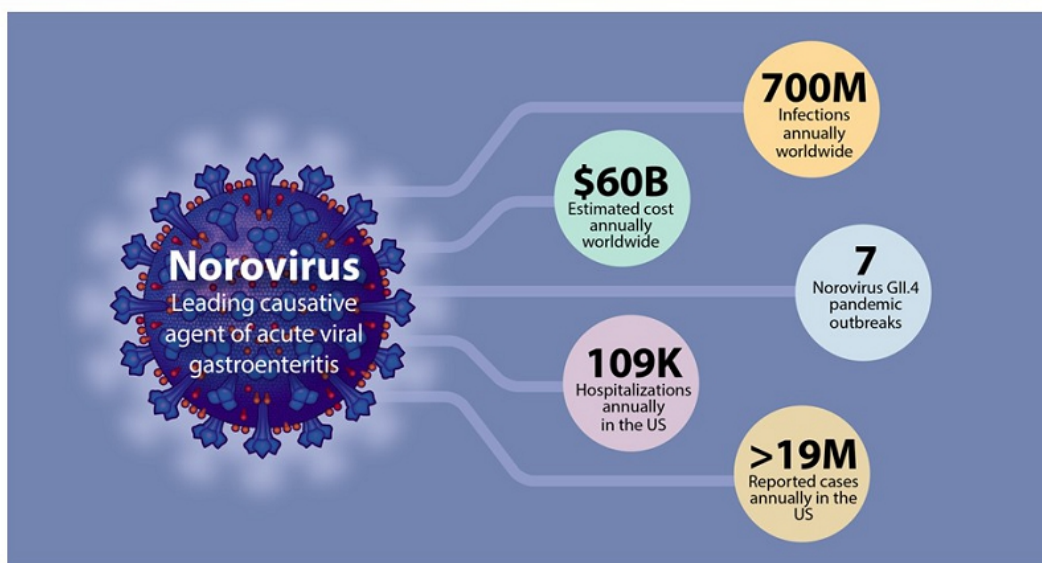
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- Favorable safety profile
- Potent, broad-spectrum activity against pandemic and seasonal strains
- High barrier to resistance
- Superior pulmonary pharmacology: high exposure and long half-life, EC<sub>50</sub> >5,000-fold by 4 days post-single administration
- Oral CC-42344: Phase 2a ongoing
- Inhaled CC-42344: Phase 1 planned in 2024

### Norovirus and Coronavirus Programs



## Norovirus Infection: No Approved Treatments or Vaccines Available



CDC: Norovirus Disease in the United States <https://www.cdc.gov/norovirus/burden.html>

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## Norovirus: Large Market with No Approved Treatments or Vaccines

- Highly contagious virus that causes symptoms of acute gastroenteritis including nausea, vomiting, stomach pain and diarrhea
- Major cause of gastrointestinal illness in closed and crowded environments including hospitals, nursing homes, childcare facilities and cruise ships
- Responsible for ~685 million infections and 200,000 deaths annually worldwide and nearly 90% of all epidemic, non-bacterial outbreaks of gastroenteritis<sup>1</sup>
  - ~200 million cases among children under five years of age and ~50,000 child deaths worldwide annually
- Estimated annual cost of \$60 billion worldwide due to direct healthcare costs and lost productivity<sup>1</sup>
- 19 million-21 million cases and 109,000 hospitalizations annually in the U.S.<sup>1</sup>

<sup>1</sup>CDC: Norovirus Disease in the United States <https://www.cdc.gov/norovirus/burden.html>

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# Norovirus and Coronavirus Development Programs Focused on Broad-spectrum Antivirals

## Clinical assets for pandemic and epidemic norovirus and coronavirus

### Oral pan-norovirus and pan-coronavirus Protease inhibitor, CDI-988

- Potential dual indications: norovirus and coronavirus
- Ongoing Phase 1 study
- Discovered by proprietary structure-based platform technology
- Potent antiviral activity against pandemic strains
- Gastrointestinal targeting activity
- Potential for both prophylactic and therapeutic treatments

## Promising Early-Stage Programs

### Replication and protease inhibitors

- Discovery ongoing
- Potent broad-spectrum activity
- Novel mechanisms of action

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## Seasoned Leadership

### Management

#### Sam Lee, Ph.D.

*Co-Chief Executive Officer & President*

25+ years of anti-infective drug discovery research experience, including HCV and influenza antivirals; played key role in early development of phosphoinositide 3-kinase (PI3K) delta inhibitor, Zydelig



#### James J. Martin, MBA, CPA

*Co-Chief Executive Officer & Chief Financial Officer*

25+ years of finance and management experience including providing financial leadership to commercial-stage, publicly traded health science companies



### Scientific Advisory Board

#### Roger Kornberg, Ph.D.

*Chairman of the Board, Chairman of the Scientific Advisory Board*

- Professor  
Stanford University School of Medicine
- Nobel Laureate

#### Michael Levitt, Ph.D.

*Member*

- Professor  
Stanford University School of Medicine
- Nobel Laureate

#### Baek Kim, Ph.D.

*Member*

- Director of Center for Drug Discovery  
Emory University

#### Bob Lehman, Ph.D.

*Member*

- Professor (Emeritus)  
Stanford University School of Medicine

#### Gary Schoolnik, M.D.

*Member*

- Professor (Emeritus)  
Stanford University School of Medicine

#### Roland Strong, Ph.D.

*Member*

- Professor  
Fred Hutchinson Cancer Research Center

#### Christophe Verlinde, Ph.D.

*Member*

- Professor (Emeritus)  
University of Washington

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## Experienced Board of Directors

### **Roger Kornberg, Ph.D.**

*Co-founder, Chairman of the Board & Chairman of the Scientific Advisory Board*

- Nobel Laureate in Chemistry
- Welch Prize – highest award granted in the field of chemistry in the U.S.
- Leopold Mayer Prize – highest award granted in the field of biomedical sciences from the French Academy of Sciences

### **Steve Rubin**

*Vice Chairman*

- EVP-Administration & Director of OPKO Health, Inc.
- Former SVP & General Counsel of IVAX Corporation; SVP & General Counsel of Telergy Inc.

### **Phillip Frost, M.D.**

*Co-founder & Director*

- Chairman & CEO of OPKO Health, Inc.
- Former Chairman of Teva Pharmaceuticals; Chairman and CEO of IVAX Corporation – sold for \$7.4 billion
- Board of Regents of Smithsonian Institution; Board of Trustees of University of Miami; Trustee of Scripps Research Institutes

### **Fred Hassan**

*Director*

- Chairman of the investment firm Caret Group; Director of global private equity firm Warburg Pincus LLC
- Former Chairman & CEO of Schering-Plough – acquired by Merck
- Former Chairman & CEO of Pharmacia Corporation; senior positions at Wyeth & Sandoz Pharmaceuticals

### **Anthony Japour, M.D.**

*Director*

- President, CEO & Director of iTolerance
- Former CEO of AdvancedDx Biological Laboratories-USA; Medical Director of ICON plc
- Former with Elite Health Medical Group specializing in infectious diseases

### **Richard C. Pfenniger, Jr.**

*Director*

- Director of OPKO Health, GP Strategies Corporation & Asensus Surgical, Inc.
- Former Chairman, CEO & President of Continucare Corporation; CEO & Vice Chairman of Whitman Education Group.
- Former COO, SVP-Legal Affairs & General Counsel of IVAX Corporation

## Expanding Intellectual Property Portfolio

### **Coronavirus**

- Issued patents in U.S. and major countries
- Pending U.S. provisional applications

### **Pandemic Influenza A**

- PB2 (influenza A inhibitor)
  - Pending applications in PCT and Taiwan
  - Pending U.S. provisional applications

### **Influenza A/B**

- Influenza A/B inhibitor
- Pending applications in U.S. and worldwide

### **Norovirus**

- Issued patents in U.S. and major countries
- Pending U.S. provisional applications

### **HCV**

#### **NS5B (NNI)**

- Issued patents in U.S.
- Pending applications in U.S. and worldwide
- Pending U.S. provisional application

## Financial Snapshot

~\$20 Million  
Market cap<sup>1</sup>

27,000  
Average 3 month  
daily share volume<sup>1</sup>

\$29.8 Million  
Cash/equivalents as of  
September 30, 2023

10.2 Million  
Common shares outstanding

10.3 Million  
Fully diluted shares

- Clean balance sheet
  - No preferred shares
  - No debt
- Cash sufficient to fund planned operations

<sup>1</sup>Yahoo Finance (November 27, 2023)

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