## 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): September 12, 2022

#### CINGULATE INC.

(Exact name of registrant as specified in its charter)

**Delaware** (State or other jurisdiction

of incorporation)

**001-40874** (Commission File Number) 86-3825535 (IRS Employer Identification No.)

1901 W. 47<sup>th</sup> Place Kansas City, KS 66205

(Address of principal executive offices) (Zip Code)

(913) 942-2300

(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 (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 exchange on which registered
Common Stock, par value \$0.0001 per share	CING	The Nasdaq Stock Market LLC
		(Nasdaq Capital Market)
Warrants, exercisable for one share of common stock	CINGW	The Nasdaq Stock Market LLC
		(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 (17 CFR §230.405) or Rule 12b-2 of the Securities Exchange Act of 1934 (17 CFR §240.12b-2). 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.  $\Box$ 

#### Item 7.01. Regulation FD Disclosure.

Cingulate Inc. updated its investor presentation to be used at investor conferences and in investor meetings. A copy of the investor presentation is furnished as Exhibit 99.1 and incorporated by reference.

#### Item 9.01. Financial Statements and Exhibits.

(d) Exhibits

Exhibit No.	Description
99.1	Investor Presentation
104	Cover Page Interactive Data File (formatted as Inline XBRL and contained in Exhibit 101)

#### **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.

#### CINGULATE INC.

Dated: September 12, 2022 By: /s/ Shane J. Schaffer

Name: Shane J. Schaffer
Title: Chief Executive Officer



## Forward-Looking Statements

This presentation contains "forward-looking statements" within the meaning of Section 27A of the Securities Act of 1933, as amended, and Section 21E of the Securities Exchange Act of 1934, as amended. These forward-looking statements include all statements, other than statements of historical fact, regarding our current views and assumptions with respect to future events regarding our business, including statements with respect to our plans, assumptions, expectations, beliefs and objectives with respect to product development, clinical studies, clinical and regulatory timelines, market opportunity, competitive position, business strategies, potential growth opportunities and other statements that are predictive in nature.

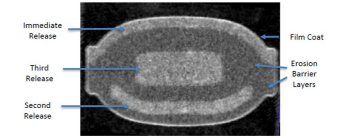
These statements are generally identified by the use of such words as "may," "could," "should," "would," "believe," "anticipate," "forecast," "estimate," "expect," "intend," "plan," "continue," "outlook," "will," "potential" and similar statements of a future or forward-looking nature. Readers are cautioned that any forward-looking information provided by us or on our behalf is not a guarantee of future performance. Actual results may differ materially from those contained in these forwardlooking statements as a result of various factors disclosed in our filings with the Securities and Exchange Commission (SEC), including the "Risk Factors" section of our Annual Report on Form 10-K for the year ended December 31, 2021. All forwardlooking statements speak only as of the date on which they are made, and we undertake no duty to update or revise any forward-looking statements, whether as a result of new information, future events or otherwise, except to the extent required



## Developing Next-Generation Medications in Billion-Dollar Markets

## Precision Timed Release™ (PTR™) Platform Unlocks the Possibility for 'True' Once-daily, Multi-dose Tablets





See the PTR™ Platform in Action



\*Symphony Data. 12-months rolling through Feb 2022



-IIS-111-0023

## Vast Pipeline of Next-Generation Medications Beyond ADHD

- Leverage PTR platform faster and with less cost in other therapeutics areas
- ✓ Market Criteria:
  - O \$1Bn+ in peak sales
  - O Next-generation mediations with significant improvement over existing therapies

#### **Identified PTR™ Platform Pipeline Opportunities**

#### **Near-Term Focus**

## **Future Therapeutic Areas**

- CTx-2103 (buspirone) Anxiety
- Insomnia
- Depression
- Bipolar Disorder Parkinson's Disease
- Cardiovascular Disorders
- Migraine
- Hypothyroidism
- Oral Oncology Medicines
- Psychosis
- Alzheimer's
- Pain (Non-Opioid)





## Targeting Treatment of ADHD - \$18Bn US Market Opportunity

Frequently diagnosed, chronic pattern interfering with functioning / development

#### ~17 Million US ADHD Patients

#### Adult ADHD

- ~11M patients in the US and growing (65% of children with ADHD become Adults with ADHD)
- 4.4% of the US adult population
- ~20% receive treatment

#### Children & Adolescents

- ~6.4M patients in the US
- 11.0% of the US under 18 population
- ~80% receive treatment

#### Societal Impact of ADHD

Estimated annual incremental costs of \$143 to \$266 billion in the United States

Earn ~ 30% less and 10% less likely to be employed

>40% higher rate of car accidents

2x greater divorce rate

2x greater incidence of accidental death

2x higher incarceration rate

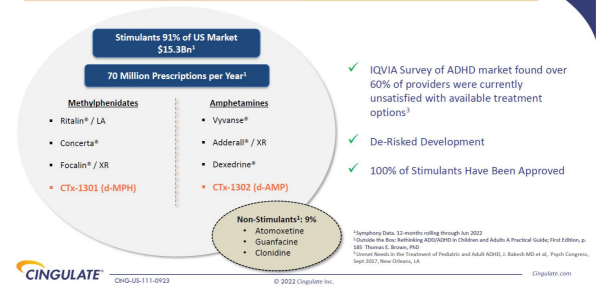


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References: https://www.cdc.gov/ncbddd/adhd/data.html
Doshi et al. J Am Acad Child Adolesc Psychiatr. 2012;51(10):990-1002.

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## \$18 Billion US ADHD Market Dominated by Stimulants



## ADHD Market Currently Dominated by 4 Stimulant Products

#### Major Unmet Medical Needs Persist

ADHD BRANDS	APPROVED	ATTRIBUTES <sup>1</sup>		UNMET NEEDS <sup>1</sup>			
		Onset	Duration (less onset)	Fast Onset of Action ≤ 30 min	Entire Active- Day Efficacy*	Minimize Crash/Rebound	Avoid Booster <sup>2</sup>
Vyvanse®	2007	2 hours	12 hours	×	×	Data Not Available	×
Adderall® XR	2001	1 ½ hours	10½ hours	×	×	Data Not Available	×
Concerta®	2000	2 hours	10 hours	×	×	Data Not Available	×
Focalin® XR	2005	30 mins	11½ hours	✓	×	Data Not Available	×



<sup>&</sup>lt;sup>1</sup> Information based upon product Package Inserts, and Summary Basis of Approvals for the approved products in chart and Ann C. Childress, Nathalie Beltran, Carl Supnet & Margaret D. Weiss (2021) Reviewing the role of emerging therapies in the ADHD armamentarium, Expert Opinion on Emerging Drugs, 26:1, 1-16.

<sup>2</sup> Symphony Data. 12-months rolling through Jun 2022



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<sup>\*</sup> Entire-active day efficacy defined as less than or equal to a 30 min onset of action with 14-16 hours of duration vs. placebo

## ADHD Market Leaders Do Not Provide "Built-In Booster"

#### Market Leaders Stop Delivery of Medication 4-5 Hours After Administration

ADHD BRANDS	ATTR	IBUTES <sup>1</sup>		RELEASE PROFILES <sup>1</sup>	
	Duration Onset (less onset)		DOSE 1 / STYLE / TIME	DOSE 2 / STYLE / TIME	DOSE 3 / STYLE /TIME
Vyvanse®	2 hours	12 hours	100% PRODRUG SUSTAINED RELEASE OVER 2 – 3 HOURS	0	0
Adderall® XR (and generics)	1½ hours	10½ hours	50% IMMEDIATE RELEASE	50% IMMEDIATE RELEASE AT HOUR 4	0
Concerta® (and generics)	2 hours	10 hours	22% IMMEDIATE RELEASE	78% SUSTAINED RELEASE OVER 4-5 HOURS	0
Focalin® XR (and generics)	30 mins	11½ hours	50% IMMEDIATE RELEASE	50% IMMEDIATE RELEASE AT HOUR 4	0

 $<sup>^{\</sup>rm 1}$  Information based upon product Package Inserts, and Summary Basis of Approvals

60%
use short-acting
'booster' dose <u>every</u>
<u>day!</u>



Source: Outside the Box: Rethinking ADD/ADHD in Children and Adults A Practical Guide; First Edition, p. 185 Thomas E. Brown, PhD

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## Recent Launches Lack Meaningful Clinical Innovation

## Niche Delivery Platforms – Designed to Fail in ADHD

ADHD BRANDS	ATTE	RIBUTES <sup>1</sup>	UNMET NEEDS				
Product	Onset	Duration	Fast Acting (≤ 30 min)	Entire Active- Day Efficacy*	Avoid Crash/Rebound	Avoid Booster	
Quillivant / Chew® XR	60 mins	8 hours	×	×	×	×	
Mydayis®	2 or 4 hrs	16+ hours	×	×	×	Potentially	
Adzenys® ER/ODT	60 mins	8-9 hours	×	×	×	×	
Cotempla® XR/ODT	60 mins	10-12 hours	×	×	×	×	
Aptensio® XR	60 mins	9 hours	×	×	×	×	
Evekeo® / ODT	60 mins	10 hours	×	×	×	×	
Dynavel® XR Oral Susp.	60 min	13 hours	×	×	×	×	
Zenzedi®	60 mins	4-5 hours	×	×	×	×	
Jornay® PM (at night)	2-hour window	10-11 hours	×	×	×	×	
Adhansia® XR	60 mins	12-13 hours	×	×	×	×	
Azstarys® (summer 2021)	Failed Endpoint	Failed Endpoint	×	×	×	×	

<sup>\*</sup> Entire-active day efficacy defined as less than or equal to a 30 min onset of action with 14-16 hours of duration vs. placebo

Information based upon product Package Inserts and Summary Sasis of Approvals and Ann C. Childress, Nathalie Beltran, Carl Supper & Margaret D. Weiss (2021) Reviewing the role of emerging therapies in the ADHD armamentarium, Expert Opinion on Emerging Drugs, 26:1, 1-16.



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# The Cingulate Solution for Patients & Providers

**ADHD** 



CING-US-111-0923

## Nine Significant Points of Differentiation

- NO ADHD product available today combines all unmet needs
- PTR™ technology affords our product candidates the following potential advantages over currently available ADHD treatments
- ✓ Provide 'entire active-day' efficacy
- Reduce abuse / diversion by eliminating booster

✓ Fast onset of action

- ✓ Significantly improved tolerability
- ✓ Eliminate need for booster/recovery dose
- ✓ Lower costs to patients, providers, and payers
- Avoid crash and rebound effect
- ✓ Ability to optimize with 8 dosage strengths
- ✓ Single-enantiomer API selection



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## CTx-1301 (d-MPH) and CTx-1302 (d-AMP)

## Ideal Design Provides Exclusive Ability to Overcome Unmet Needs

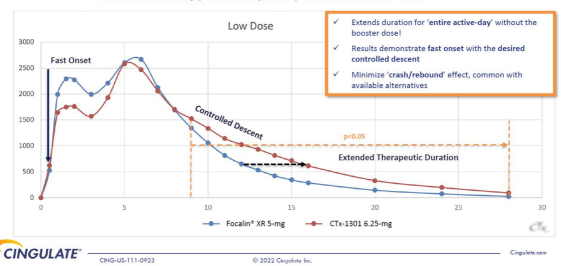
CINGULATE	TARGET	ATTRIBUTES			
	Onset	Duration	DOSE 1 / STYLE / TIME	DOSE 2 / STYLE / TIME	DOSE 3 / STYLE /TIME
CTx-1301 (d-MPH)	30 mins	Up to 16 hours	35% IMMEDIATE RELEASE	45% SUSTAINED RELEASE OVER 90 MINUTES AT HOUR 3	20% IMMEDIATE RELEASE AT HOUR 7
CTx-1302 (d-AMP)	30 mins	Up to 16 hours	45% IMMEDIATE RELEASE	35% SUSTAINED RELEASE OVER 90 MINUTES AT HOUR 3	20% IMMEDIATE RELEASE AT HOUR 7

CINGULATE	TARGET	ATTRIBUTES	UNMET NEEDS			
	Onset	Duration	Fast Acting (≤ 30 min)	Entire Active- Day Efficacy	Avoid Crash/Rebound	Avoid Booster
CTx-1301 (d-MPH)	30 mins	Up to 16 hours	✓	✓	✓	✓
CTx-1301 (d-AMP)	30 mins	Up to 16 hours	✓	✓	✓	✓

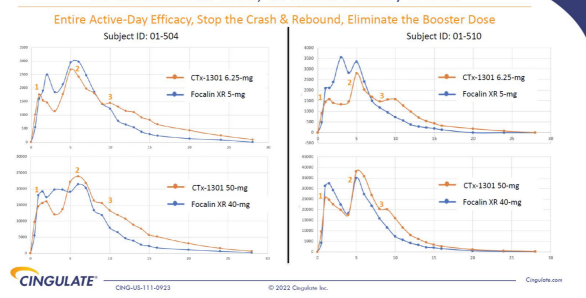


## CTx-1301 Clinical Phase 2 Study Results

#### Plasma dexmethylphenidate (dMPH) Concentration vs Time



## At the Individual Level, Tri-modal Delivery is Clear



## CTx-1301 Demonstrated Significantly Lower Adverse Events

#### 28.6% reduction in TEAE's related to CTx-1301 versus Focalin XR (14.3% difference)

	Focalin XR 5 mg (n=41)	CTx-1301 6.25 mg (n=39)	Focalin XR 40 mg (n=43)	CTx-1301 50 mg (n=42)	All CTx-1301 (n=42)	All Focalin XR (n=44)
atients with at least one						
Treatment Emergent Adverse Events	7 (17.1%)	4 (10.3%)	22 (51.2%)	14 (33.3%)	17 (40.5%)	25 (56.8%
Mild	7 (17.1%)	4 (10.3%)	20 (46.5%)	14 (33%)	17 (40.5%)	23 (52.3%
Moderate	0	0	2 (4.7%)	0	0	2 (4.5%)
Severe	0	0	0	0	0	0
TEAE Related to Study Drug	5 (12.2%)	3 (7.7%)	20 (46.5%)	13 (31.0%)	15 (35.7%)	22 (50.0%
AE Leading to Study or Drug Withdrawal	1 (2.4%)	0	1 (2.3%)	0	0	2 (4.5%)

There were no serious adverse events.

Source: CSR CTx-1301-001 Listing 16.2.7.1



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#### Commercialization to Drive Revenue

#### Changing dynamics in ADHD commercial landscape

- · Ability to dominate share of voice
  - o Concerta, Adderall XR, Focalin XR are all offpatent with no promotion
  - Vyvanse loss of exclusivity ~August 2023
- New entrants lack major promotional efforts, field forces, and revenue

#### **Maximize Access for Patients and Providers**

- Clinical, Practical, and Societal Story:
  - Efficacy and Tolerability
  - One versus Two Prescriptions
  - Abuse & Diversion
- Rebates & Net to Plan Cost
  - PBM's driven by rebate guarantees to payers; estimated >\$2B last year\*
  - ADHD is a high brand utilization market with high-cost generics at 55-90% of branded drug cost\*

#### Cingulate's Comprehensive Commercial Model

- > Branded product of choice ~ Patients, Providers, & Payers
- > Strategic partnership to maximize market access, distribution, promotion across all channels
  - Psychiatry / Neurology & Pediatrics / Family Practice encompass 84% of ADHD market\*
  - Maximize and retain NPV to Cingulate



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\*Symphony Data. 12-months rolling through Jun 2022

## Exclusivity: IP, Agreements, and Trade Secrets

#### Intellectual property rights expected to provide exclusivity through 2035 at a minimum

OralogiK™ Erosion Barrier Layer



- Five (5) patents granted (US & Global) expiry dates ranging from 2031 to 2035,
- One (1) OralogiK™ patent pending (US, Europe)
- Three (3) Cingulate product specific patents under prosecution with USPTO and global entities
  - Pharmacokinetics
- Pharmacodynamics
- Trimodal release of API
- Formulation, Precise Timing, Ratio of API

#### **Exclusivity agreements**



- Compression technology exclusivity for branded Cingulate products
- Significant modifications and exclusive process technologies incorporated



Methods, tools, processes, designs, and equipment trade secrets







Develop...

Shape market acceptance, and...

Prepare to commercialize next-generation drug candidates...

Where currently prescribed standard-of-care treatments result in suboptimal outcomes for all stakeholders

Achievement Drives Shareholder and Team Member Value



