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): February 27, 2023

TONIX PHARMACEUTICALS HOLDING CORP.

(Exact name of registrant as specified in its charter)

Nevada (State or Other Jurisdiction of Incorporation) 001-36019 (Commission File Number) 26-1434750 (IRS Employer Identification No.)

26 Main Street, Chatham, New Jersey 07928 (Address of principal executive offices) (Zip Code)

Registrant's telephone number, including area code: (862) 904-8182

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	TNXP	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 \Box

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.

Tonix Pharmaceuticals Holding Corp. (the "Company") updated its investor presentation, which is used to conduct meetings with investors, stockholders and analysts and at investor conferences, and which the Company intends to place on its website, which may contain nonpublic information. A copy of the presentation is filed as Exhibit 99.01 hereto and incorporated herein by reference.

The information in this Item 7.01 of this Current Report on Form 8-K, including Exhibit 99.01 attached hereto, shall not be deemed "filed" for purposes of Section 18 of the United States Securities Exchange Act of 1934 (the "Exchange Act") or otherwise subject to the liabilities of that section, nor shall they be deemed incorporated by reference in any filing under the United States Securities Act of 1933 or the Exchange Act, except as shall be expressly set forth by specific reference in such a filing.

Item 9.01 Financial Statements and Exhibits.

(d)	Exhibit	
_	No.	Description.
_	99.01	Corporate Presentation by the Company for February 2023
	104	Cover Page Interactive Data File (embedded within the Inline XBRL document)

SIGNATURE

Pursuant to the requirement 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.

TONIX PHARMACEUTICALS HOLDING CORP.

By: <u>/s/ Bradley Saenger</u> Bradley Saenger Chief Financial Officer

Exhibit 99.01

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Cautionary Note on Forward-Looking Statements

Certain statements in this presentation regarding strategic plans, expectations and objectives for future operations or results are "forward-looking statements" as defined by the Private Securities Litigation Reform Act of 1995. These statements may be identified by the use of forward-looking words such as "anticipate," "believe," "forecast," "estimate" and "intend," among others. These forward-looking statements are based on Tonix's current expectations and actual results could differ materially. There are a number of factors that could cause actual events to differ materially from those indicated by such forward-looking statements. These factors include, but are not limited to, the risks related to failure to obtain FDA clearances or approvals and noncompliance with FDA regulations; delays and uncertainties caused by the global COVID-19 pandemic; risks related to the timing and progress of clinical development of our product candidates; our need for additional financing; uncertainties of patent protection and litigation; uncertainties of government or third party payor reimbursement; limited research and development efforts and dependence upon third parties; and substantial competition. As with any pharmaceutical under development. there are significant risks in the development, regulatory approval and commercialization of new products. The forward-looking statements in this presentation are made as of the date of this presentation, even if subsequently made available by Tonix on its website or otherwise. Tonix does not undertake an obligation to update or revise any forward-looking statement, except as required by law. Investors should read the risk factors set forth in the Annual Report on Form 10-K for the year ended December 31, 2021, as filed with the Securities and Exchange Commission (the "SEC") on March 14, 2022, and periodic reports and current reports filed with the SEC on or after the date thereof. All of Tonix's forward-looking statements are expressly qualified by all such risk factors and other cautionary statements.

Who We Are

OUR MISSION



Tonix Pharmaceuticals is committed to improving population health by inventing and developing innovative therapies and vaccines, through broad in-house capabilities and creative collaborations, to help address important unmet needs.

OUR VISION

Tonix strives to be a leader in providing **novel drug therapies and** vaccines to improve population health around the world.

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Investment Highlights



DIVERSE PIPELINE

Tonix's core focus is on **central nervous system** disorders, but we also target unmet needs across multiple therapeutic areas including **immunology**, **infectious disease** and **rare disease**.



IN-HOUSE CAPABILITIES

Investment in domestic, **in-house**, **R&D** and **manufacturing** to accelerate development timelines and improve the ability to respond to pandemics.



STRATEGIC PARTNERSHIPS

Partnering strategically with other **biotech companies**, **world-class academic and non-profit** research organizations to bring innovative therapeutics to market faster.



FINANCIAL POSITION

Tonix had approximately \$120 M in cash and cash equivalents as of 12/31/22. Tonix has no debt.

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TONIX

Pipeline: Key Programs

andidates*	Indication	Status/Next Milestone	
TNX-102 SL ¹	Fibromyalgia (FM) Long COVID (PASC ²)	Mid-Phase 3 - >50% enrolled Phase 2 - enrolling	
TNX-1300 ³	Cocaine Intoxication - FDA Breakthrough Designation	Mid-Phase 2, Targeted 2Q 2023 Start	
TNX-1900 ⁴	Migraine, Craniofacial Pain and Binge Eating Disorder	Phase 2 - enrolling	() N
TNX-601 ER	Depression, PTSD, Neurocognitive Dysfunction from Steroids	Phase 2, Targeted 1Q 2023 Start ⁶	Man
TNX-16007	Depression, PTSD and ADHD	Preclinical	and and
TNX-29008	Prader-Willi Syndrome - FDA Orphan Drug Designation	Preclinical	
TNX-15009	Organ Transplant Rejection/ Autoimmune Conditions	Phase 1, Targeted 2Q 2023 Start	10
TNX-170010	Gastric and colorectal cancers	Preclinical	
TNX-801 ¹¹	Smallpox and mpox vaccine	Phase 1, Targeted 2H 2023 Start	
TNX-1850 ¹²	COVID-19 Vaccine (horsepox-based live virus vaccine)	Preclinical	
TNX-230013	COVID-19 Vaccine	Preclinical	
TNX-360014	COVID-19 Therapeutic Platform (fully human monoclonal antibodies)	Preclinical	
TNX-370015	COVID-19 Vaccine (zinc nanoparticle mRNA technology)	Preclinical	
TNX-380016	COVID-19 Therapeutic/Preventative (humanized monoclonal antibodies)	Preclinical	
e HCI sublingual tablets /ID-19. ccaine esterase) was lici ense agreement with Str stigator-initiated IND ha development was comp rrma; license agreemen	new drugs or biologics and have not been approved for any indication. Is also in development for Agitation in Alzheimer's Disease (AAD) and Alcohol Use Disorder (AUD). Both indications are ensed from Columbia University. anford University, IND cleared for the prevention of migraine indication; Planned Binge Eating Disorder study is expected to as been completed in the U.S. using TNX-1900; Phase 2 for the prevention of migraine headache expected to start 1Q 2023 level outside of the U.S. Phase 2 expected to start 1Q 2023 twith Wayne State University I institute of Headh and Medical Research (Inserm)	"Live attenuated vaccine based on horsepox virus "Live attenuated vaccine based on horsepox virus vector, expressed SARB-CoV2 spike protein. TWX1850 is based on the BA.2 variant spike protein. "Live attenuated vaccine based on bovine parainfluenza (BPI) virus "Fully human monocional antibody generated from COVID-19 convaliseout patients "COVID vaccine based on mRNA in zinc nanoparticle (ZNP) formulation with CD40, molecular trigger "Humanized monocional antibody generated from mice immunized with SARB-CoV2 spike protein	т¢і

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ll of Tonix's pro NX-102 SL (cy ase 2 ready.

(4.100) (double-mutant cocaine estrate) was licensed from Columbia University. guided from Trigenina; license agreement with Stanford University; IND cleared for the prevention of migraine indication; Planned Binge Eating Disorder study is expected to versignation initiated. be investigator initiated. W Phase 2 that under an investigator-initiated IND has been completed in the U.S. using TNX-1900; Phase 2 for the prevention of migraine headache expected to start 1Q 2023 Acquired from TRimaran Pharma; license agreement with Wayne State University Co-exclusive Iclonese agreement with French National Institute of Health and Medical Research (Inserm) ant-CD4U, humanited monochoral antibody Recombinant trefo factor2 (TFR2) based protein; licensed from Columbia University © 2023 Tonix Pharmaceuticals Holding (

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Five Late-Stage CNS Programs to be in the Clinic by 1H'23 Three Studies Enrolling Now

ENROLLING

- In Phase 3:
 - TNX-102 SL for fibromyalgia (>50% enrolled)
- In Phase 2:
 - TNX-102 SL for fibromyalgia-type Long COVID
 - TNX-1900 for migraine headache (new mechanism for US patients)

Entering Phase 2 in 1H23

- TNX-601 ER for major depressive disorder (new mechanism for US patients)
- TNX-1300 for cocaine intoxication (breakthrough therapy designation)

Potential Pivotal Study

Potential Pivotal Study

Potential Pivotal Study Potential Pivotal Study



CNS PORTFOLIO

TNX-102 SL*

1Not approved for any indication

Cyclobenzaprine (Protectic[®]) Pipeline in a Product

A unique, sublingual formulation of cyclobenzaprine designed to optimize delivery and absorption

Potent binding and antagonist activities at the serotonin-5-HT2A, α 1-adrenergic, histaminergic-H1, and muscarinic-M1 receptors to facilitate restorative sleep

Innovative and proprietary PROTECTIC[®] Rapid drug exposure following nighttime administration

Differentiators:

Relative to Oral Cyclobenzaprine

- Lower daytime exposure
- Avoids first-pass metabolism
- Reduces risk of pharmacological interference from major metabolite

Relative to Standard of Care

· Potential for better tolerability while maintaining efficacy

Patents Issued

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Fibromyalgia

Long COVID

Status: Phase 2

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Status: Mid-Phase 3

>50% enrolled

One positive Phase 3 study (RELIEF) completed

· Phase 2 study (PREVAIL) is currently enrolling

Next Steps: Trial enrollment is in process

One Phase 2 study (AtEase) completed

Posttraumatic Stress Disorder (PTSD)

Two Phase 3 studies (HONOR, RECOVERY) conducted

Second Phase 3 study (RALLY) missed primary endpoint

Next Steps: Interim analysis results expected 2Q 2023

Confirmatory Phase 3 study (RESILIENT) is currently enrolling

TNX-102 SL*: Fibromyalgia Cyclobenzaprine Protectic[®] Sublingual Tablets

PROFILE

Fibromyalgia (FM) is a chronic pain disorder resulting from amplified sensory and pain signaling within the CNS

- Afflicts an estimated 6-12 million adults in the U.S., approximately 90% of whom are women¹
- Symptoms include chronic widespread pain, nonrestorative sleep, fatigue, and cognitive dysfunction
- Patients struggle with daily activities, have impaired quality of life, and frequently are disabled
- Physicians and patients report common dissatisfaction with currently marketed products



When the check engine light malfunctions, the light is on even though the car is not malfunctioning

Patents Issued

¹American Chronic Pain Association (www.theacpa.org, 2019)

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DEVELOPMENT PROGRAM

Market Entry: Fibromyalgia

Additional Indications: Long COVID, PTSD, Agitation in Alzheimer's, Alcohol Use Disorder

Status: One Positive Phase 3 study RELIEF completed

Second Phase 3 study RALLY missed primary endpoint

Confirmatory Phase 3 study RESILIENT is currently enrolling

Next Steps: Interim analysis results expected 2Q 2023

*TNX-102 SL has not been approved for any indication



CNS PORTFOLIC

NS PORTFOLIO

Phase 3 RESILIENT Study Design

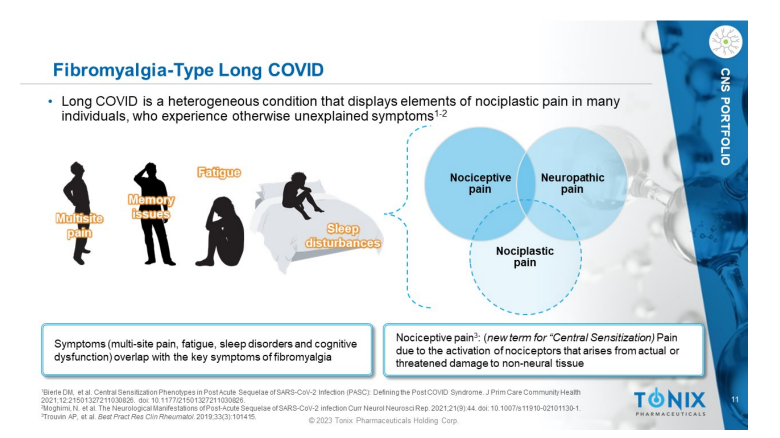
General study characteristics:

- · Randomized, double-blind, placebo-controlled study in fibromyalgia
- · U.S. sites only, expected to enroll approximately 470 patients
- One unblinded interim analysis based on 50% of randomized participants

Primary Endpoint:

- Daily diary pain severity score change from baseline to Week 14 (TNX-102 SL vs. placebo)
 - · Weekly averages of the daily numerical rating scale scores
 - · Analyzed by mixed model repeated measures with multiple imputation (MMRM with MI)





TNX-102 SL*: Long COVID (PASC) Cyclobenzaprine Protectic[®] Sublingual Tablets

PROFILE

- Occurs in approximately 13% of recovered COVID-19 patients¹
- As many as 40% of Long COVID patients experience multi-site pain, a hallmark of fibromyalgia^{2,3}



Patents Issued

September 1, 2022- CDC - https://www.cdc.gov/coronavirus/2019-ncov/long-term-effects/index.html ?Harris, H, et al. Tonix data on file. 2022 3TriNetX Analytics

DEVELOPMENT PROGRAM

Market Entry: Fibromyalgia-Type Long COVID (PASC)

Additional Indications: Fibromyalgia, PTSD, Agitation in Alzheimer's, Alcohol Use Disorder

Status: Phase 2 study PREVAIL is currently enrolling

Next Steps: PREVAIL trial enrollment is in process

ClinicalTrials.gov Identifier: NCT05472090 A Phase 2 Study to Evaluate the Efficacy and Safety of TNX-102 SL in Patients With Multi-Site Pain Associated With Post-Acute Sequelae of SARS-CoV-2 Infection (PREVAIL)

*TNX-102 SL has not been approved for any indication.



12

CNS PORTFOLIO

Phase 2 Long COVID Study Design (PREVAIL)

Study characteristics:

- Randomized, double-blind, placebo-controlled study of TNX-102 SL in fibromyalgia-type Long COVID
- · U.S. sites only, expected to enroll approximately 470 patients
- · One unblinded interim analysis based on 50% of randomized participants

Primary Endpoint:

- Daily diary pain severity score change from baseline to Week 14 (TNX-102 SL vs. placebo)
 - · Weekly averages of the daily numerical rating scale scores
 - · Analyzed by mixed model repeated measures with multiple imputation (MMRM with MI)

TNX-102 SL once-daily at bedtime 5.6 mg (2 x 2.8 mg tablets)^{*}

Placebo once-daily at bedtime

14 weeks —

Two week run in at 2.8 mg dose at bedtime, followed by 12 weeks at 5.6 mg dose

ClinicalTrials.gov Identifier: NCT05472090 "A Phase 2 Study to Evaluate the Efficacy and Safety of TNX-102 SL in Patients With Multi-Site Pain Associated With Post-Acute Sequelae of SARS-CoV-2 Infection (PREVAIL)"

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TNX-601 ER*: Depression Tianeptine Hemioxalate Extended-Release Tablets (39.4 mg)

PROFILE

- · A novel, oral, extended-release once-daily tablet
- Treatment effect of tianeptine sodium immediate release t.i.d. in depression is well-established
- Tianeptine stimulates neurogenesis in animal models
- Indirectly modulates the glutamatergic system
- Does not interact with AMPA, NMDA or Kainate receptors¹

Differentiators:

Relative to Tianeptine IR:

Once daily dosing

Relative to traditional anti-depressants:

- Unique mechanism of action
- Tianeptine sodium IR has similar efficacy but fewer side effects than traditional anti-depressants

Patents Issued

¹AMPA=α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid; NMDA=N-methyl-D-aspartate

DEVELOPMENT PROGRAM

Market Entry: Major Depressive Disorder

Additional Indications: PTSD, Neurocognitive Disorder From Corticosteroids

Status: Phase 2 ready

Next Steps: Initiate a Phase 2 potentially pivotal study 1Q 2023

- Double-blind, placebo-controlled, parallel-group, randomized
- 6-week treatment period
- Expected to enroll approximately 300 patients across 30 sites in the US

*TNX-601 ER has not been approved for any indication



NS PORTFOLIO

CNS PORTFOLIO

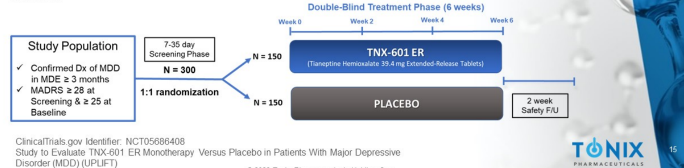
TNX-601 ER - Phase 2 UPLIFT Study Design

General study characteristics:

- Randomized, double-blind, placebo-controlled study in Major Depressive Disorder
- · Parallel design (two arms-treatment (tianeptine hemioxalate 39.4 mg) and placebo)
- ~30 U.S. sites only, expected to enroll approximately 300 patients
- One unblinded interim analysis based on 50% of randomized participants expected 4Q'23

Primary Endpoint:

 Mean change in the Montgomery Åsberg Depression Rating Scale week 6, change from baseline

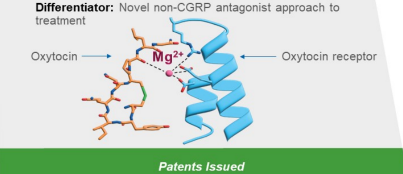


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TNX-1900*: Migraine Intranasal Potentiated Oxytocin (OT) with Magnesium

PROFILE

- Intranasal OT has potential utility in treating migraine¹
- Magnesium is known to potentiate the binding of OT to its receptor^{2,3}
- One billion individuals worldwide suffer from migraines



DEVELOPMENT PROGRAM

Market Entry: Chronic Migraine

Additional Indications: Acute Migraine, Craniofacial Pain, Insulin Resistance, Binge Eating Disorder

> Status: Phase 2 study PREVENTION is currently enrolling⁴

Next Steps: Interim analysis results expected 4Q 2023

Investigator initiated Phase 2 trial in binge eating disorder 1Q 2023

CNS PORTFOLIO

CNS PORTFOLIO

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*TNX-1900 has not been approved for any indication. CGRP = calcitonin ge

Tzabazis A, et al. Oxytocin and Migraine Headache. Headache. 2017 May;57 Suppl 2:64-75. doi: 10.1111/head.13082. PMID: 28485846.
*Antoni FA, Chadio SE, Essential role of magnesium in oxytocin-receptor affinity and ligand specificity. Biochem J. 1999 Jan 15;257(2):611-4. doi: 10.1042/bj2570611. PMID: 2539090; PMCID: PMC1135623.
*Meyerowitz, 10., *et al.* The oxytocin signating complex reveals a molecular switch for cation dependence. *Nat Struct Mol Biol* (2022), (https://doi.org/10.1038/s41594-022-00728-4)
*A Phase 2 trial under an investigator-initiated IND has been completed in the U.S. using TNX-1900
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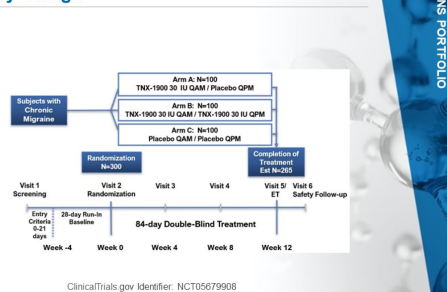
Phase 2 PREVENTION Study Design

General study characteristics:

- Randomized, double-blind, placebo-controlled study (three arms– two treatment regimens and one placebo) in chronic migraine
- U.S. sites only, expected to enroll approximately 300 patients
- One unblinded interim analysis based on 50% of randomized participants expected 4Q'23

Primary Endpoint:

 Mean change in the number of migraine headache days between the 28-day Run-In phase and the last 28-days of the Treatment phase (TNX-1900 vs. placebo)



A Study to Evaluate the Efficacy and Safety of TNX-1900 in Patients With Chronic Migraine (PREVENTION)

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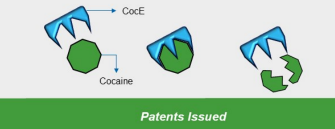
TNX-1300*: Cocaine Intoxication Cocaine Esterase (CocE)

PROFILE

Cocaine is the main cause for drug-related ED visits¹ CocE is a recombinant protein that degrades cocaine in the bloodstream

- Rapidly reverses physiologic effects of cocaine
- Drops plasma exposure by 90% in 2 minutes

Differentiators: Rapidly metabolizes cocaine in the bloodstream; no other product currently on the market for this indication



DEVELOPMENT PROGRAM

Market Entry: Cocaine Intoxication

Status: Mid-Phase 2

Next Steps: Initiate new Phase 2 trial 2Q 2023 pending FDA agreement

CNS PORTFOLIO

18

- Single-blind, placebo (+ usual care) controlled, randomized, potentially pivotal study
- Expected to enroll approximately 60 emergency department patients at sites in the US

FDA Breakthrough Therapy Designation

Awarded Cooperative Agreement Grant from National Institute on Drug Abuse (NIDA)

¹Havakuk O et al. *J Am Coll Cardiol*. 2017;70:101-113. ED = emergency department.

© 2023 Tonix Pharmaceuticals Holding Corp. *TNX-1300 has not been approved for any indication.

RARE DISEASE: KEY CANDIDATES

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RARE DISEASE PORTFOLIO

20

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TNX-2900*: Hyperphagia in Prader-Willi Syndrome Intranasal Potentiated Oxytocin (OT) with Magnesium

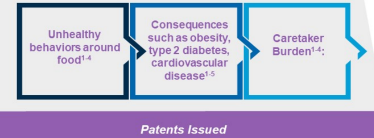
PROFILE

Prader-Willi Syndrome is the most common genetic cause of life-threatening childhood obesity

 Rare disease occurring in 1 in 10,000 to 1 in 30,000 births

Differentiator: No approved therapeutic currently on the market for hyperphagia in PWS

Dangers of PWS Hyperphagia:



 'Miller JL, et al. Am J Med Genet A. 2011;155A(5):1040-1049.

 'Butler MG, et al. Genet Med. 2017;19(6):635-642.

 'Butler MG. NORD. Updated 2018. Accessed May 25, 2022. https://rarediseases.org/rare-diseases/prader-willi-syndrome/

 'Prader-Will Syndrome Association USA. Accessed May 25, 2022. https://www.pvsausa.org/what-is-prader-willi-syndrome/

 'Prader-Will Syndrome Association USA. Accessed May 25, 2022. https://www.pvsausa.org/what-is-prader-willi-syndrome/

 'Muscogiuri G, et al. J Endocrinol Invest. 2021;44(10):2057-2070.

DEVELOPMENT PROGRAM

Market Entry: Hyperphagia in Prader-Willi Syndrome

Additional Indications: Rare Hyperphagia Conditions

Status: Pre-IND

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Next Steps: IND preparation

*TNX-2900 is in the pre-IND stage of development and has not been approved for any indication.



TNX-1500*



Next Generation α -CD40 Ligand (CD40L) Antibody

The CD40-CD40L pathway is a pivotal immune system modulator and a well-established and promising treatment target

Differentiators: Expected to deliver efficacy without compromising safety

First Generation: Development halted due to thromboembolic (TE) complications—blood clots—traced to Fc gamma receptor (Fc γ R)

Second Generation: Eliminated the Fc γ R TE complication but potency and half life was reduced, limiting utility

Third Generation (TNX-1500): Re-engineered to better modulate the binding of $Fc\gamma R$ while preserving FcRn function.

*TNX-1500 is in the pre-IND stage of development and has not been approved for any indication. Patents filed.

Prevention of Allograft Rejection

Status: Preclinical

 Collaborations ongoing with Mass General Hospital on heart and kidney transplantation in non-human primates

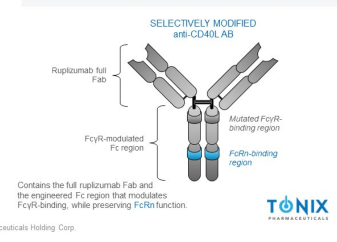
Next Steps: Initiate Phase 1 study 2Q 2023

Autoimmune Diseases

Status: Potential future indications include:

Sjögren's Syndrome, Systemic Lupus Erythematosus

These indications require large studies, but represent large target markets



Other anti-CD40L Monoclonal Antibodies in Development

UCB (Co-developed with Biogen) - Systemic Lupus Erythematosus (SLE)

- Phase 3 Trial Currently Enrolling (NCT04294667)
- Topline results expected 1H 2024¹
- Dapirolizumab pegol (pegylated Fab)

Horizon (Agreed to be acquired by Amgen) - Sjögren's Syndrome (SjS)

- Two Positive Phase 2 studies reported^{2,3} .
- Dazodalibep (tn03 fusion protein)

Sanofi - Sjögren's Syndrome (SjS), Multiple Sclerosis (MS), Systemic Lupus Erythematosus (SLE)

- Phase 2 Trial Currently Enrolling in SiS (NCT04572841) and SLE (NCT05039840)
- Active Phase 2 Trial in Relapsing MS (NCT04879628)
- SAR441344 (Fc-modified)

Eledon – Amyotrophic Lateral Sclerosis (ALS) and Kidney Transplant

- Phase 2 Trial Completed in ALS (NCT04322149)
- Phase 1/2 Trial Currently Enrolling in Kidney Transplant (NCT05027906)
- Tegoprubart, f.k.a. AT-1501 (Fc-modified) •

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¹https://www.ucb.com/our-science/pipeline ²https://ir.horizontherapeutics.com/news-releases/news-release-details/horizon-therapeutics-plc-announces-phase-2-trial-evaluating ³https://ir.horizontherapeutics.com/news-releases/news-release-details/horizon-therapeutics-plc-announces-phase-2-trial-evaluating-0 © 2023 Tonix Pharmaceuticals Holding Corp.

TNX-1700*: Gastric and Colorectal Cancers **Recombinant Trefoil Factor 2 (rTFF2) Fusion Protein**

Potential New Cancer Treatment

- TNX-1700 (rTFF2) has effects on cancer by altering the tumor micro-environment
- Mechanism of action: suppresses myeloid-derived suppressor cells and activates anti-cancer CD8+T cells
- Potential synergy with anti-PD-1 or anti-PD-L1 monoclonal • antibodies (mAbs)

Preclinical Evidence for Inhibiting Growth of **Cancer Cells**

 Data showed that TFF2-CTP augmented the efficacy of mAb anti-PD-1 therapy. Anti-PD-1 in combination with TFF2-CTP showed greater anti-tumor activity in PD-L1overexpressing mice

Licensed from Columbia University

Developing in partnership under sponsored research agreement

Patents Filed

Market Entry: Immuno-oncology, combination therapy with PD1 blockers for gastric and colorectal cancer

Status: Preclinical

Next Steps: Animal studies ongoing

Differentiator: No product yet identified consistently augments PD1 effects on cold tumors

*TNX-1700 is in the pre-IND stage of development and has not been approved for any indication.

IMUNOLOGY PORTFOLIO

IMMUNOLOGY PORTFOLIO

24



TNX-801 & TNX-1850*



Recombinant Pox Vaccine (RPV) Platform Using Live Virus Technology

Differentiators:

- Live virus vaccines are the most established vaccine technology
 - Starting with Edward Jenner's smallpox vaccine, the first vaccine, which eradicated smallpox
 - Prevents forward transmission
 - Effective in eliciting durable or long-term immunity
- · Economical to manufacture at scale
 - Low dose because replication amplifies dose in vivo
 - Single shot administration
- Standard refrigeration required for shipping and storage

*TNX-801 and TNX-1850 are in the pre-IND stage of development and has not been approved for any indication. Patents filed. Noyce RS, et al. Construction of an infectious horsepox virus vaccine from chemically synthesized DNA fragments. PLoS One. 2018 Jan 19;13(1):e018453. 'Brennan, Z: Endpoints March 2, 2022 (https://endpts.com/weaker-omicron-variant-is-great-news-for-the-world-but-bad-newsfor a nort of entitied values (https://endpts.com/weaker-omicron-values-for-the-world-but-bad-newsfor a nort o

Mpox and Smallpox Vaccine

Status: Preclinical

• TNX-801 is a cloned version of horsepox¹ (without any insert) purified from cell culture

Next Steps: Developing GMP manufacturing; Initiate Phase 1 Trial 2H 2023

COVID-19 Vaccine

Status: Preclinical

- First version TNX-1800 encodes spike protein from SARS-CoV-2, Wuhan strain
- Planned new version TNX-1850 encode spike protein from SARS-CoV-2 BA.2 strain²

Next Steps: Developing TNX-1850 (BA.2) version



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TNX-3600*: COVID-19 Therapeutic/Preventative Fully Human Monoclonal Antibody

PROFILE

- · Fully human monoclonal antibodies
- Generated from SARS-CoV-2+ asymptomatic individuals or COVID-19 convalescent patients
- Potential to be scaled up quickly and combined with other monoclonal antibodies
- · Collaboration with Columbia University

TXX-3800⁴ Mouse hybridoma approach¹ The manual of the sponse of the

DEVELOPMENT PROGRAM

Market Entry: COVID-19 treatment and prophylaxis in immuno-compromised individuals

Status: Preclinical

Next Steps: Study inhibition of SARS CoV-2 variants in tissue culture; initiate animal studies in 1H 2023

Differentiators: Potential to decrease response time to newly identified COVID-19 variants, relative to generating murine mAbs followed by humanization NFECTIOUS DISEASE PORTFOLIO

27

INFECTIOUS

DISEASE PORTFOLIO

28

*TNX-3600 is in the pre-IND stage of development and has not been approved for any indication

¹Lu R-M, Hwang Y-C, Liu JJ, et al. Development of the rapeutic antibodies for the treatment of diseases. J Biomed Sci. 2020;27(1):1. doi:10.1186/s12929-019-0592-z ²TNX-3600 and TNX-3800 are the designations for a series of monoclonal antibodies; each is in the pre-IND stage of development and has not been approved for any indication

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TNX-3800*: COVID-19 Therapeutic/Preventative Humanized Mouse Monoclonal Antibodies

PROFILE

- · Humanized monoclonal antibodies
- · Generated from mice immunized with SARS-CoV-2 spike protein
- · Exclusive license from Curia Global, Inc.

Differentiators: To date, EUA-approved products have been derived from the blood of COVID-convalescent patients or a humanized mouse^{1,2}

Relative to fully humanized mAbs:

- Murine mAbs discovered by Curia and licensed by Tonix represent a potential new approach to treating SARS-CoV-2 infection
- Murine mAbs have the potential to neutralize a broader spectrum of SARS-CoV-2 variants and <u>may be more difficult to evade</u> as we face expanding variant pool from both convergent and divergent evolution³

Patents Filed

DEVELOPMENT PROGRAM

Market Entry: COVID-19 treatment and prophylaxis in immuno-compromised individuals

Status: Preclinical

Next Steps: Study inhibition of SARS CoV-2 variants in tissue culture; initiate animal studies in 1H 2023

*TNX-3800 is in the pre-IND stage of development and has not been approved for any indication.

¹Hansen J et al. Science. 2020 Aug 21;369(6506):1010-1014. Doi: 10.1126/science.abd0827 ²Asdaq, S.M.B. et al. A Patent Review on the Therapeutic Application of Monoclonal Antibodies in COVID-19. Int. J. Mol. Sci. 2021, 22, 11953. <u>https://doi.org/10.3390/iims222111953</u> ³Callaway, E. oct 28 2022. Nature (News), COVID Variants oury is making winter surges hard to predict. Descendants of Omicron are proliferating worldwide — and the same mutations are coming up again and again. www.nature.com/articles/d41586-022-03445-6 © 2023 Tonix Pharmaceuticals Holding Corp.

Additional Infectious Disease Therapeutics in Development

TNX-2300*: Live Virus Vaccine Based on Bovine Parainfluenza (BPI) Virus Market Entry: COVID-19 Vaccine Status: Preclinical Next Steps: Animal studies with Kansas State University (KSU) to test the effect of co-expression of CD40-ligand to stimulate T cell immunity VFECTIOUS DISEASE PORTFOLIO

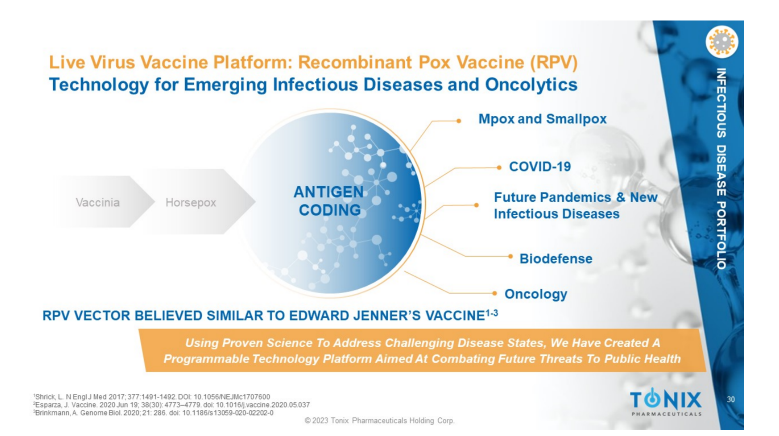
 TNX-3700*: Zinc Nanoparticle (ZNP) Formulation for mRNA Vaccines

 Market Entry: Booster for COVID-19 Vaccines

 Status: Preclinical

 Next Steps: Research at KSU on CoV-2 spike based vaccine in tissue culture and animals; initiate animal studies in 1H 2023

*TNX-2300 and TNX-3700 are in the pre-IND stage of development and have not been approved for any indication.



Internal Development & Manufacturing Capabilities

Infectious Disease R&D Center (RDC) – Frederick, MD

- Function: Accelerated development of vaccines and antiviral drugs against COVID-19, its variants and other infectious diseases
- Description: ~48,000 square feet, BSL-2 with some areas designated BSL-3
- Status: Operational

Advanced Development Center (ADC) - North Dartmouth, MA

- · Function: Development and clinical scale manufacturing of biologics
- Description: ~45,000 square feet, BSL-2
- Status: Operational

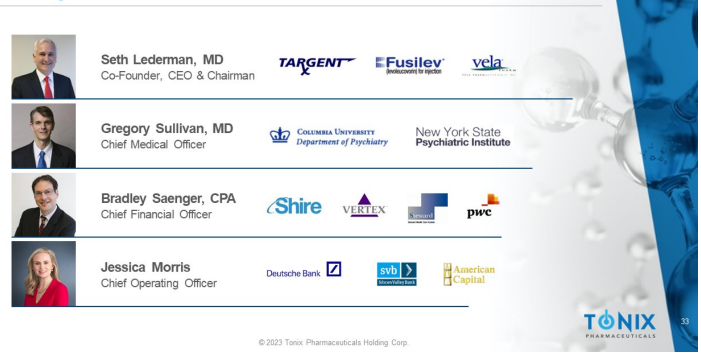
Commercial Manufacturing Center (CMC) - Hamilton, MT

- Function: Phase 3 and Commercial scale manufacturing of biologics
- Description: ~44-acre green field site, planned BSL-2
- Status: Planning for site enabling work in 2023





Management Team



Milestones: Recently Completed and Upcoming*

- ₽ 2nd Quarter 2022 Phase 3 RESILIENT study start of TNX-102 SL for the management of fibromyalgia
- 3rd Quarter 2022 Phase 2 PREVAIL study start of TNX-102 SL for the treatment of Long COVID
- 1st Quarter 2023 Phase 2 study start of TNX-1900 for the treatment of migraine

Expected Data

- 2nd Quarter 2023 Interim Analysis results of Phase 3 RESILIENT study of TNX-102 SL for fibromyalgia
- □ 4th Quarter 2023 Interim Analysis results of Phase 2 PREVENTION study of TNX-1900 for chronic migraine
- □ 4th Quarter 2023 Interim Analysis results of Phase 2 study of TNX-601 for major depressive disorder

Expected Clinical Trial Initiations

- □ 1st Quarter 2023 Phase 2 study start of TNX-601 ER for the treatment of major depressive disorder
- 2nd Quarter 2023 Phase 1 study start of TNX-1500 for prevention of allograft rejection
- □ 2nd Quarter 2023 Phase 2 study start of TNX-1300 for the treatment of cocaine intoxication
- □ 2nd Half 2023 Phase 1 study start of TNX-801 for prevention of mpox and smallpox

* We cannot predict whether the global COVID-19 pandemic will impact the timing of these milestones.

THANK YOU