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 6, 2022

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 September 2022
	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



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.

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What We Do



OUR MISSION

ADVANCING THE SCIENCE AND UNDERSTANDING OF DISEASES by developing **innovative therapies** that improve **population health** by focusing on **unmet needs** in patient care

OUR STRATEGY

Using our integrated development engine, we advance innovative programs across multiple therapeutic areas into the clinic while maximizing asset potential



OUR UNIQUE CAPABILTIIES

With our diverse and deep pipeline, broad team of experienced professionals covering all facets of drug development, as well as in-house R&D and manufacturing abilities, Tonix can respond quickly to **new emerging diseases** and threats

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Milestones: Recently Completed and Upcoming*

1st Quarter 2022	Topline data from Phase 3 RALLY study of TNX-102 SL for the management of fibromyalgia
🖬 2 nd 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
Expected Data	Interim analysis results of Phase 3 RESILIENT study of TNX-102 SL in fibromyalgia
□ 1 st Half 2023	Interim analysis results of Phase 2 PREVAIL study of TNX-102 SL in Long COVID
Expected Clinical Tr	rial Initiations Phase 2 study start of TNX-102 SL for the treatment of PTSD in Kenya
4 th Quarter 2022	Phase 2 study start of TNX-1900 for the treatment of migraine
1st Quarter 2023	Phase 2 study start of TNX-601 ER for the treatment of major depressive disorder
1 st Quarter 2023	Phase 2 study start of TNX-1300 for the treatment of cocaine intoxication
1st Half 2023	Phase 1 study start of TNX-1500 for prevention of allograft rejection
□ 1 st Half 2023	Phase 1 study start of TNX-801 for prevention of monkeypox and smallpox in Kenya

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

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ANDIDATES*	INDICATION	STATUS / NEXT MILESTONE
TNX-102 SL ¹	Fibromyalgia (FM) Posttraumatic Stress Disorder (PTSD) Long COVID (PASC ²)	Mid-Phase 3 Phase 2, Targeted 3Q 2022 Start Phase 2
TNX-13003	Cocaine Intoxication FDA Breakthrough Designation	Mid-Phase 2, Targeted 1Q 2023 Start
TNX-1900 ⁴	Migraine, Craniofacial Pain and Binge Eating Disorder	Phase 2, Targeted 4Q 2022 Start ⁵
TNX-601 ER	Depression, PTSD, Neurocognitive Dysfunction from Steroids	Phase 2, Targeted 1Q 2023 Start
TNX-16007	Depression, PTSD and ADHD	Preclinical
benzaprine HCI sublingual ae of COVID-19. mutant cocalne esterase) i emina; license agreement i er an investigator-initiated he pre-IND stage in the U.3	ational new drugs or biologics and have not been approved for any indication. tablets) is also in development for Agtablen in Alzheimer's Disease (AAD) and Alcohol Us was licensed fram Columbia University. with Stanford University. IND cleared for the prevention of migraine indication; Planned Bir IND has been completed in the U.S. using TWX-1900; Phase 2 for the prevention of migra 5; a Phase 1 trial for formulation development was completed outside of the U.S; Phase 2 ement with Warve State University.	nge Eating Disorder study is expected to be investigator i ine headache expected to start 40,2022

Pipeline **Rare Disease Portfolio**

CANDIDATES*	INDICATION	STATUS / NEXT MILESTONE
TNX-29001	Prader-Willi Syndrome FDA Orphan Drug Designation	Preclinical

#

Pipeline

Immunology and Immuno-Oncology portfolio

	CANDIDATES*	INDICATION	STATUS / NEXT MILESTONE	
	TNX-29001	Prader-Willi Syndrome FDA Orphan Drug Designation	Preclinical	1
pelin	icense agreement with French Nat	nar new drags or biologics and have not been approved for any indication. ional Institute of Health and Medical Research (Inserm)		R
	CANDIDATES*	INDICATION	STATUS / NEXT MILESTONE	
R	CANDIDATES*	INDICATION Organ Transplant Rejection/ Autoimmune Conditions		- 1

Pipeline Infectious Disease Portfolio

CANDIDATES*	INDICATION	STATUS / NEXT MILESTONE
TNX-8011	Smallpox and monkeypox vaccine	Phase 1, Targeted 1H 2023 Star
TNX-1850 ²	COVID-19 Vaccine (horsepox-based live virus vaccine)	Phase 1, Targeted 2H 2023 Sta
TNX-23003	COVID-19 Vaccine	Preclinical
TNX-36004	COVID-19 Therapeutic Platform (monoclonal antibodies)	Preclinical
TNX-37005	COVID-19 Vaccine (zinc nanoparticle mRNA technology)	Preclinical
tenuated vaccine based on horsepox v	irus vector, expressed SARS-CoV-2 spike protein. TNX-1850 is based on the BA.2 variant spi ainfluenza (BPI) virus	ke protein.



TNX-102 SL*: Fibromyalgia Cyclobenzaprine Protectic® Sublingual tablets

PROFILE

A unique formulation of cyclobenzaprine designed to optimize delivery and absorption

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

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

Clinical trial program designed to examine treatment of core Fibromyalgia symptoms

Patents Issued

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 NS PORTFOLIO

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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.

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TNX-102 SL: Fibromyalgia Program Update

Phase 3 Study, RESILIENT, will compare TNX-102 SL 5.6 mg and placebo

- First patient enrolled in April 2022
- Interim Analysis results expected 2Q 2023
- · Parallel design, double-blind, randomized placebo-controlled study, all U.S. sites
- · Primary endpoint is pain at Week 14 analyzed by MMRM with MI
- Projecting adverse event-related discontinuations to decrease towards rates in RELIEF and PTSD Studies

Phase 3 Study, RALLY, comparison of TNX-102 SL 5.6 mg and placebo

- As expected from interim analysis results published in July 2021, RALLY Study missed primary endpoint
- Unexpected ~80% increase in adverse event-related discontinuations in both drug and placebo arms
- Multiple imputation approach on 'Missing Data' attenuated statistical significance of efficacy endpoints'
- TNX-102 SL was generally well tolerated with overall adverse event profile comparable to prior studies; no new safety signals observed

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

PROFILE

Long COVID or Post-acute Sequelae of COVID-19 (PASC¹)

- Symptoms can include fatigue, sleep disorders, pain, fevers, shortness of breath, cognitive impairment described as "brain fog", gastrointestinal symptoms, anxiety, and depression²
- Can persist for months and can range in severity from mild to incapacitating
- Occurs in 30% of recovered COVID-19 patients
- Typically associated with moderate or severe COVID-19, Long COVID can occur after mild COVID-19 or even after asymptomatic SARS-CoV-2 infection

DEVELOPMENT PROGRAM

Market Entry: Long COVID (PASC)

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

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Status: Phase 2 study PREVAIL is currently enrolling

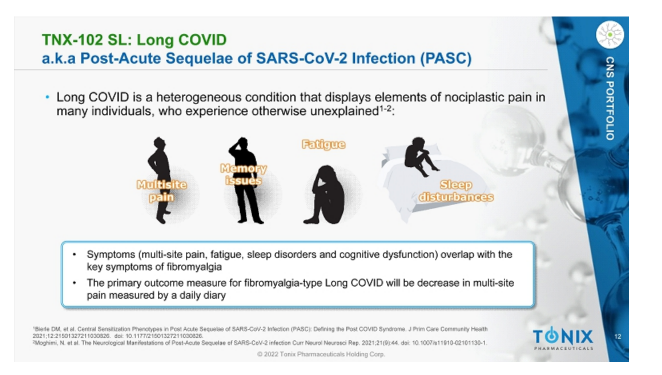
Next Steps: Interim analysis results expected 1H 2023

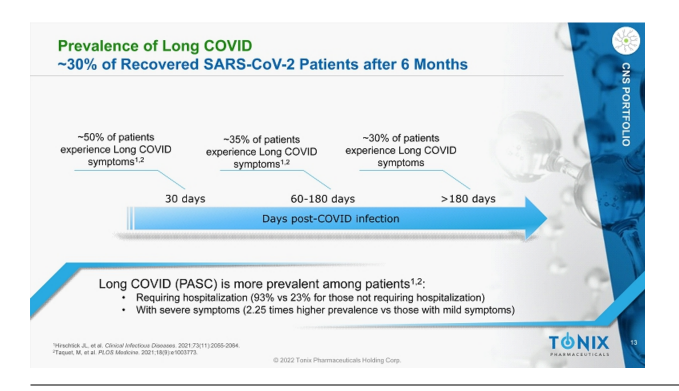
*TNX-102 SL has not been approved for any indication

Patents Issued

Feb. 24, 2021 - White House COVID-19 Response Team press briefing; Feb 25, 2021 - policy brief from the World Health Organization on long COVID "Nationalian, Ani, et al. "Post-acute COVID-19 syndrome." Nature Medicine (2021); 1-15.

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Rate of Opioid Use in Long COVID Patients Potential Health Concern

In only a few days some people can develop a physical dependence and addiction to opioids¹⁻²

The USA Department of Labor estimates that 1 in 4 patients prescribed opioids long term will struggle with opioid addiction adding to the already growing opioid crisis¹⁻²

¹Shah, A, et al. MMWR Morb Mortal Willy Rep. 2017;66:265–269.
²U.S. Department of Labor

IS PORTFOLIO 100% Percent of Patients taking Opioids 90% 80% 70% 51.1% 50.1% 60% Legend: Long COVID symptoms 50% 39.0% No multi-site pain 80 40% X Multi-site pain only 30% Multi-site pain and fatigue 20% Multi-site pain and insomnia 10% Muti-site pain, fatigue, and insomnia 0% Source: Harris, H, et al. Tonix data on file. 2022.; TriNetX Analytics τοΝΙΧ © 2022 Tonix Pharmaceuticals Holding Corp

TNX 102 SL*: Posttraumatic Stress disorder (PTSD) Cyclobenzaprine Protectic® Sublingual Tablets

PROFILE

PTSD is a serious chronic psychiatric illness

 Defined as maladaptive prolonged stress response which occurs after experiencing severely injurious traumatic event(s)

Affects approximately 12 million Americans adults^{1,2}

Large unmet clinical need and limited effective therapies available

 Advances in pharmacological treatments beyond the currently approved SSRIs (e.g., Zoloft® (sertraline), Paxil® (paroxetine)) are needed³

Patents Issued

DEVELOPMENT PROGRAM

Market Entry: PTSD

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

Status: One Phase 2 study (AtEase) completed

Two Phase 3 studies (HONOR, RECOVERY) conducted

> Next Steps: 3Q 2022 Initiate Phase 2 Trial in Kenya

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

³Cain, C. K., et al. Targeting memory processes with drugs to preve or cure PTSD. Expert Opin Investig Drugs. 2012; 21(9), 1323-1350

¹Coldstein RB, et al. The epidemiology of DBM-5 postraumatic stress disorder in the United States: results from the National Epidemiologic Survey on Acohol and Related Conditions-III. Soc Psychiatry Psychiatr Epidemiol. 2016;51(b):1137-1148, Paintzak RH, et al. Prevalence and Asia Icomotifyed of full and partial posttrauratic States disorder in the United States: results from Vave 2 of the National Epidemiologic Survey on Alcohol and Related Conditions. J Anelety Disord. 2011;25(3):456-465 (2) 2022 Tonix Pharmaceuticals Holding Corp.

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

PROFILE

Cocaine is the main cause for drug-related $\ensuremath{\mathsf{ED}}\xspace$ visits^1

Cocaine use can cause irreversible structural damage to the heart and accelerate cardiovascular disease²

 In one survey of 94 long-term cocaine users, 71% had some form of cardiovascular disease³

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

Patents Issued

¹Havakuk O et al. J Am Coll Cambol. 2017;70:101-113. ²PHilips K et al. Am J Cardiovasc Drugs. 2009;9:177-196. ³Maceira AM et al. J Cardiovasc Magn Reson. 2014;16:26. ED = emergency department.

DEVELOPMENT PROGRAM

Market Entry: Cocaine Intoxication

Status: Mid-Phase 2

Next Steps: Initiate a new Phase 2 single-blind, placebo-controlled, randomized, potentially pivotal study in 1Q 2023, pending FDA agreement.

FDA Breakthrough Therapy Designation

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

*TNX-1300 has not been approved for any indication © 2022 Tonix Pharmaceuticals Holding Corp.

TNX-601 ER*: Depression Tianeptine Hemioxalate Extended-Release Tablets

PROFILE

A novel, oral, extended-release once-daily tablet

Mechanistically different from traditional monoaminergic treatments for depression

Indirectly modulates the glutamatergic system

 No direct binding to NMDA, AMPA, or kainate receptors

Treatment effect of tianeptine in depression is well-established



Market Entry: Major Depressive Disorder

Additional Indications: PTSD, Neurocognitive Disorder From Corticosteroids CNS PORTFOLIO

CNS PORTFOLIO

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Status: pre-IND

Next Steps: 1Q 2023 Initiate Phase 2 Trial

Patents Issued

*TNX-601 ER is in the pre-IND stage of development and has not been approved for any indication.

Additional Indications: Acute Migraine,

Status: Clinical - IND cleared for

prevention of migraine headache4

Next Steps: 4Q 2022 Initiate Phase

2 Trial and Investigator Initiated Phase 2 Trial in Binge Eating

Craniofacial Pain, Insulin Resistance,

AMPA=a-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid; MAOI=monoamine oxidase inhibitors; NMDA=N-methyl-D-aspartate

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

PROFILE

DEVELOPMENT PROGRAM Market Entry: Chronic Migraine

Binge Eating Disorder

Disorder

Intranasal OT has potential utility in treating migraine¹

- · Intranasal OT reaches the trigeminal ganglion
- Preclinical evidence of OT blocking CGRP release and suppressing pain
- Association of low OT levels during and preceding migraine episodes
- Novel non-CGRP antagonist approach to treatment

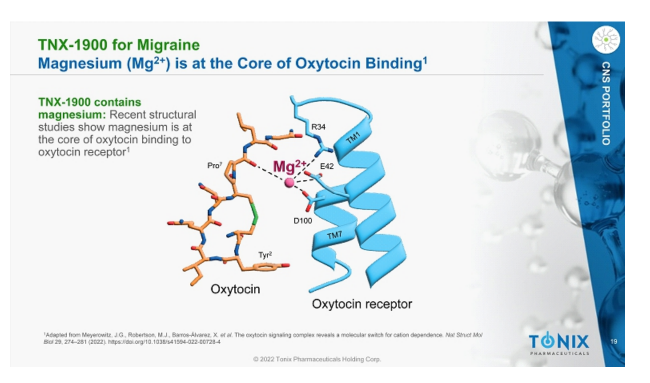
Magnesium is known to potentiate the binding of OT to its receptor^{2,3}

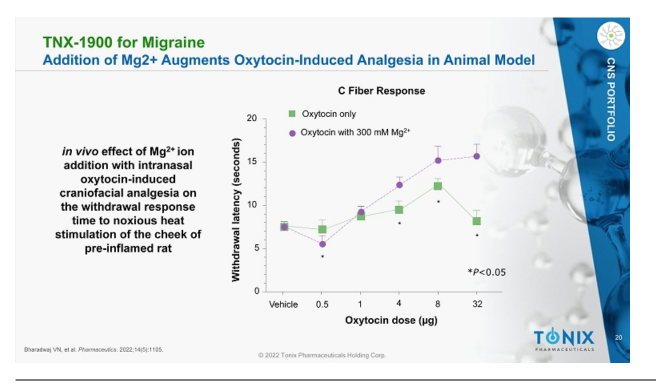
One billion individuals worldwide suffer from migraines

Patents Issued

*TNX-1900 has not been approved for any indication. CGRP = calcitonin gene-related peptide.

17 zabazis A, et al. Oxylocin and Migraine Headache. Headache. 2017 May;57 Suppl 2:64-75. doi: 10.1111/head.13082. PMID: 2848546. Antoni FA, Chadio SE, Essential role of magnesium in oxylocin-receptor affinity and Igand specificity. Biochem J. 1899 Jan 15:257(2);611-4. doi: 10.1042/bj2570611. PMID: 2539090; PMCID: PMC1135







TNX-2900*: 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

Symptoms include lack of suckling as infants, poor muscle strength, and constant hunger (hyperphagia)

- In animal models, OT has improved suckling and suppressed hunger
 - Tonix's patented potentiated OT formulation is believed to increase specificity for OT receptors relative to off-target vasopressin receptors

Patents Issued

DEVELOPMENT PROGRAM

Market Entry: Prader-Willi Syndrome

Additional Indications: Rare Hyperphagia Conditions

Status: Preclinical, granted orphan drug designation by FDA

Next Steps: pre-IND Meeting to seek agreement on development plans ARE DISEASE PORTFOLIO

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

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TNX-1500*: Prevention of Allograft Rejection 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

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

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

Third Generation (TNX-1500): Re-engineered to better modulate the binding of Fc γR while preserving FcRn function

Expected to deliver efficacy without compromising safety

Status: Preclinical; collaborations ongoing with Mass General Hospital on heart and kidney transplantation in non-human primates

Next Steps: 1H 2023 Initiate Phase 1 Study

Patents Filed

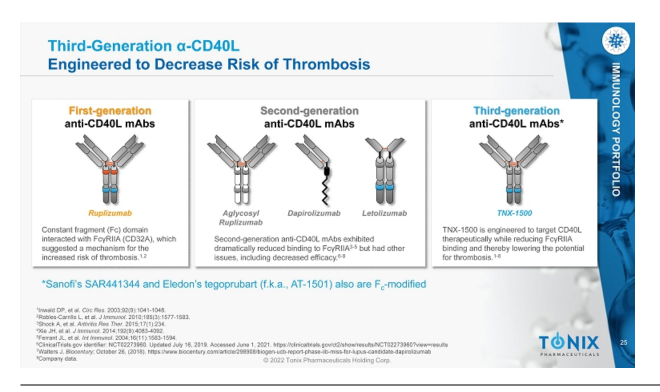
'Camilleri B, et al. Exp Clin Transplant. 2016;14(5):471-483.

Rupiizumab full Fab GryR-modulated Fc region CryR-binding region GryR-binding region Contains the full rupiizumab Fab and the engineered Fc region that modulates fcyR-binding, while preserving FcRn function.

SELECTIVELY MODIFIED

anti-CD40LAB

MMUNOLOGY PORTFOLIO



Development and Regulatory Strategy IMMUNOLOGY PORTFOLIO 1st Indication – Kidney allotransplantation (human to human) Replacement for nephrotoxic CNI's (calcineurin inhibitors, e.g. Prograf® (tacrolimus)¹, Neoral® (cyclosporin)² - Similar development path to the successful development of BMS's Nulojix® (belatacept)3, CTLA-4/Ig biologic - Clinical development may combine with Nulojix or Rapamune® (rapamycin/sirolimus)4 2nd Indication – Heart or kidney xenotransplant (pig to human) CD40L:CD40 blockade considered essential The engineered pig organ is also considered a biologic 3rd Indication –Lou Gehrig's Disease, or ALS⁵ Animal models show strong activity; competitor Eledon (ELDN) is pursuing ALS as primary indication · 4th Indication (and beyond) - Autoimmune disease (e.g., Systemic Lupus Erythematosus, Rheumatoid Arthritis, Progressive Systemic Sclerosis) These indications require large studies; SLE and RA would represent very large target markets ¹http://www.accessdata.idda.gov/drugsatfda_itocs/label/2009/050708s027,050709s0211bl.pdf ²http://www.novartis.us/istest/www.novartis.us/Ties/heoral.pdf ²http://backageinserts.bms.com/pib/_nulojuc.pdf ⁴https://backageinserts.bms.com/pib/_nulojuc.pdf ⁴https://backageinserts.bms.com/_nulojuc.pdf ⁴https://backageinserts.bms.com/_nuloiuc.pdf ⁴https:// τΰΝΙΧ © 2022 Tonix Pharmaceuticals Holding Corp



TNX-801: Monkeypox and Smallpox Vaccine Live Virus Platform Development Program

APPLICATION OF LIVE VIRUS PLATFORM

- TNX-801 is a cloned version of horsepox¹ (without any insert) purified from cell culture
- In addition to being a potential addition to the U.S. Strategic National Stockpile, TNX-801serves as the basis for the RPV/horsepox platform

ANIMAL TESTING OF TNX-801 WITH SOUTHERN RESEARCH INSTITUTE

 Non-human primate monkeypox challenge testing: positive data reported in 1Q 2020²

DEVELOPED IN COLLABORATION WITH UNIVERSITY OF ALBERTA

 Proprietary synthetic biology approach and vector system

Patents Filed

DEVELOPMENT PROGRAM

Market Entry: Monkeypox and Smallpox Vaccine

Status: Preclinical, Pre-IND

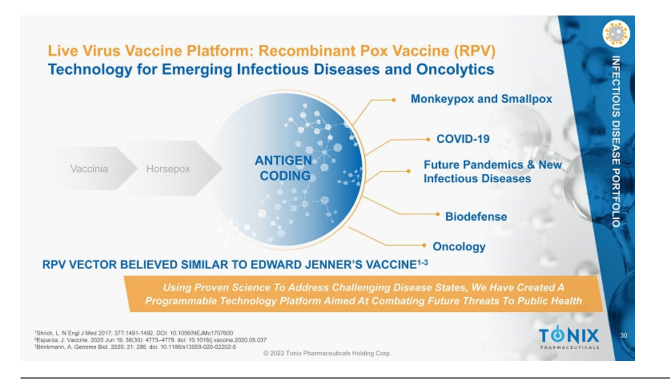
Next Steps: Developing GMP manufacturing for TNX-801; initiate Phase 1 Trial, 1H 2023 in Kenya FECTIOUS DISEASE PORTFOLIO

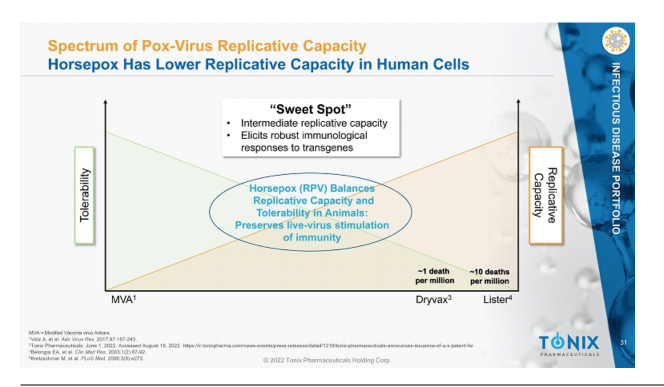
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*TNX-801 is in the pre-IND stage of development and has not

been approved for any indication.

Noyce RS, et al. Construction of an infectious horsepox virus vaccine from chemically synthesized DNA fragments. PLoS One. 2018 Jan 19;13(1):e0188453. Noyce, RS, et al. Synthetic Chimeric Horsepox Virus (scHPXV) Vaccinstion Protects Macaques from Monkeypox? Presented as a poster at the American Society of Microbiology BioThreats Canterence - January 29, 2020, Arlington, VA. (https://content.equisolve.net/tonixpharma/media/0023ac2714tb/6f/3204/Erd 165a121, pdf) © 2022 Tonix Pharmaceuticals Holding Corp.





Vaccinia and Horsepox Induce a Skin Reaction Called a "Take" Described by Dr. Edward Jenner NFECTIOUS DISEASE PORTFOLIO Biomarker of protection - Smallpox was eradicated using this marker - Revaccination indicated for recipients without "take" Take² Measure of T cell immunity No need for blood draws or complex laboratory studies No other functional T cell assay is approved or in clinical use for vaccination Intradermal vaccination¹ *Example of major cutaneous reaction, or "take," resulting from a replication-competent live-virus vaccine with intradermal delivery, indicating successful vaccination1.2 'Fulginiti VA, et al. Cliv Infect Dis. 2009;37(2):241-250. ²Centers for Disease Control and Prevention. Accessed April 15, 2020. https://phil.cdo.gov/Details.aspx?pid=3276 τΰΝΙΧ 32 © 2022 Tonix Pharmaceuticals Holding Corp

TNX-1850*: COVID-19 Vaccine Live Virus Platform Development Program

APPLICATION OF LIVE VIRUS PLATFORM

- 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 strain¹

ANIMAL TESTING OF TNX-1800 WITH SOUTHERN RESEARCH INSTITUTE

- Non-human primate immune response: positive results reported in 4Q 2020
- Non-human primate CoV-2 challenge testing: positive data reported in 1Q 2021

DEVELOPED IN COLLABORATION WITH UNIVERSITY OF ALBERTA

· Proprietary synthetic biology approach and vector system

Patents Filed

DEVELOPMENT PROGRAM

Market Entry: COVID-19 Vaccine

Additional Indications: Future Pandemic, Infectious Disease, Smallpox, Cancer

Status: Preclinical

Next Steps: Developing TNX-1850 (BA.2) version; initiate Phase 1 Trial, 2H 2023

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*TNX-1850 is in the pre-IND stage of development and has not been approved for any indication.

Brennan, Z. Endpoints March 2, 2022 (https://endpts.com/weaker-omicron-variant-is-great-news-for-the-world-but-bad-news-for-covid-related-clinical-trials/)

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Live Virus Platform: What Makes TNX-1850 Different from mRNA Vaccines

CRITERIA	mRNA VACCINES	TNX-1850
Number of shots	Two	One
Duration	6 months	Years / decades
Boosters	Recommended	Likely not required
Protection from variants	Decreased	Expected
Forward transmission	Unknown for variants	Likely prevents
Biomarker	None	Yes – "Take"
Manufacturing	Complex	Conventional
Glass-sparing packaging	No	Yes
Shipping and storage	Cold chain	Standard refrigeration
Protection from smallpox	No	Yes
* Characterizations of TNX-1850 shown in table	represent expectations.	
		чот

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TNX-2300*: COVID-19 Vaccine Live Virus Vaccine Based on Bovine Parainfluenza (BPI) Virus

LIVE VIRUS VACCINE¹⁻⁵

- Previously has been shown to be an effective antigen delivery vector in humans, notably well tolerated in infants and children
- Vector is well suited for mucosal immunization using a nasal atomizer, but it can also be delivered parenterally

ANIMAL TESTING OF TNX-2300 ONGOING

- Non-human primate immune response: positive results reported in 4Q 2020
- Non-human primate CoV-2 challenge testing: positive data reported in 1Q 2021

DEVELOPED IN COLLABORATION WITH KANSAS STATE UNIVERSITY (KSU)

 Uses a novel live attenuated vaccine vector platform, BPI, and the CD40-ligand to stimulate T cell immunity

Patents Filed

*TNX-2300 is in the pre-IND stage of development and has not

Next Steps: Animal studies with KSU to test

also known as CD154 or 5c8 antigen, to

the effect of co-expression of the CD40-ligand,

FECTIOUS DISEASE PORTFOLIO

NFECTIOUS DISEASE PORTFOLIO

been approved for any indication.

 ¹Halle, AA et al. J Gen. Virology (2003) 84:2153–2162; ²Halle, AA et al. J Virology (2000) 74 (24): 11626–11635; ³Karron RA et al. J Inf Dis (1995) 171: 1107-14; ⁴Karron RA et al. J Inf Dis (1995) 171: 1107-14; ⁴Karron RA et al. J Virology (2001) 75(10): 4594–4603

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TNX-3600*: COVID-19 Therapeutics Fully Human Monoclonal Antibody Platform

PROFILE

Collaboration with Columbia University

Human monoclonal antibodies (mAbs) generated from COVID-19 convalescent patients

Potential monotherapies

 Plan to seek indication similar to current EUA therapeutic mAbs for treating individuals with mild-to-moderate COVID-19 who are at high risk for progression to severe disease

Potential combination therapy with other antibodies Combination therapies for other anti-CoV-2 monoclonal

 Combination therapies for other anti-cov-z monocional antibodies are believed to have reduced the emergence of drug resistant viral strains

Given the unpredictable trajectory of the SARS-CoV-2 virus and new variants¹, we seek to contribute to a broad set of monoclonal antibodies from a variety of patients, that can be scaled up quickly and potentially combined with other monoclonal antibodies.

Waltz, E. Nature. "Does the World Need an Omicron Vaccine?" 28 Jan 2022 https://www.nature.com/articles/d41588-022-00199-z

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DEVELOPMENT PROGRAM

DEVELOPMENT PROGRAM

Infectious Diseases

Status: Preclinical

Market Entry: COVID-19 Vaccine

stimulate T cell immunity.

Additional Indications: Future Pandemic,

Market Entry: COVID-19 Therapeutic

Additional Indications: Symptomatic COVID in patients with risk factors for poor outcome

Status: Preclinical

Next Steps: Study inhibition of SARS CoV-2 variants in tissue culture; 2Q 2022 Initiate Animal Studies

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

TNX-3700*: COVID-19 Vaccine Zinc Nanoparticle (ZNP) Formulation for mRNA Vaccines

PROFILE

Collaboration with Kansas State University

ZNP technology is a potential replacement for the Lipid Nanoparticle (LNP) technology of current mRNA vaccines

Potential improved stability

- Plan to seek initial indications as booster, similar to the current EUA and FDA approved mRNA vaccines
- · Improved stability would facilitate shipping and storage

Addresses limitations in current mRNA vaccines

which require ultra-cold storage and shipping
 Stability issues limit use in less developed countries

Patents Filed

Market Entry: Booster for COVID-19 Vaccines

DEVELOPMENT PROGRAM

Additional Indications: COVID-19 vaccine for naïve individuals

Status: Preclinical

Next Steps: Research at K-State on CoV-2 spike based vaccine in tissue culture and animals; 2Q 2022 Initiate Animal Studies FECTIOUS DISEASE PORTFOLIC

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*TNX-3700 is in the pre-IND stage of development and has not bee approved for any indication.

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Live Virus RPV Platform & COVID-19 Vaccine Internal Development & Manufacturing Capabilities

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

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

Advanced Development Center (ADC) - North Dartmouth, MA

- <u>Function</u>: Development and clinical scale manufacturing of live-virus vaccines
- Description: ~45,000 square feet, BSL-2
- <u>Status</u>: Partially operational as of 2Q 2022

Commercial Manufacturing Center (CMC) - Hamilton, MT

- <u>Function</u>: Phase 3 and Commercial scale manufacturing of live-virus vaccines
- · Description: ~44 acre green field site, planned BSL-2
- · Status: Planning for site enabling work in 2022

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American Pandemic Preparedness Plan (AP3)

"Platforms" – Foundation of Pandemic Response

- Key element of AP3 from White House Office of Science and Technology Policy or OSTP^{1,2}
 - 100 days to human trials
 - Technologies that do not require sterile injection

TNX-801/TNX-1850 (live virus RPV) platform addresses OSTP requirements^{1,2}

 Our goal is to be able to test new live virus vaccines against novel pathogens within the 100 days of obtaining sequence FECTIOUS DISEASE PORTFOLIO

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- RDC is equipped to make new vaccines
- ADC will be equipped to make clinical trial material
- · CMC is planned to make commercial scale material

¹ Sept 3, 2021 (https://www.whitehouse.gov/wp-content/uploads/2021/09/American-Pandemic-Preparedness-Transforming-Our-Capabilities-Final-For-Web.pdf) ² Sept 3, 2021 (https://www.whitehouse.gov/briefing-room/statements-releases/2021/09/03/fact-sheet-biden-administration-to-transform-capabilities-for-pandemic-preparedness/) © 2022 Tanix Pharmaceuticals Holding Corp.





Milestones: Recently Completed and Upcoming*

1st Quarter 2022	Topline data from Phase 3 RALLY study of TNX-102 SL for the management of fibromyalgi	a
Y 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) (m
Expected Data	Interim analysis results of Phase 3 RESILIENT study of TNX-102 SL in fibromyalgia	Martin
□ 1 st Half 2023	Interim analysis results of Phase 2 PREVAIL study of TNX-102 SL in Long COVID	d ras
Expected Clinical T	rial Initiations	
3rd Quarter 2022	Phase 2 study start of TNX-102 SL for the treatment of PTSD in Kenya	
4 th Quarter 2022	Phase 2 study start of TNX-1900 for the treatment of migraine	
□ 1 st Quarter 2023	Phase 2 study start of TNX-601 ER for the treatment of major depressive disorder	C 100
□ 1 st Quarter 2023	Phase 2 study start of TNX-1300 for the treatment of cocaine intoxication	0.27
1st Half 2023	Phase 1 study start of TNX-1500 for prevention of allograft rejection	
1st Half 2023	Phase 1 study start of TNX-801 for prevention of monkeypox and smallpox in Kenya	a. 12
"We cannot predict whether the	global COVID-19 pandemic will impact the timing of these milestones.	τόνιχ
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