

**Corporate Presentation** 

Focus on: Tonmya<sup>™\*</sup> (TNX-102 SL) in Development for the Management of Fibromyalgia

**April 2024** 

NASDAQ: TNXP

\*Tonmya is conditionally accepted by FDA as the tradename for TNX-102 SL for the management of fibromyalgia



## **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; risks related to the failure to successfully market any of our products; 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, 2023, as filed with the Securities and Exchange Commission (the "SEC") on April 1, 2024, 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

Tonix is committed to developing and marketing therapeutics to treat pain, neurologic, psychiatric and addiction conditions through our *central* nervous system portfolio and within other areas of high unmet need, including immunology, infectious disease, and rare disease

## With a Focus on:

Filing a New Drug Application (NDA) with the US Food and Drug
Administration (FDA) for Tonmya™ (TNX-102 SL) for the management
of Fibromyalgia



## CNS-Focused Biopharma with Preclinical, Clinical and Commercial Stage Products



## Tonmya<sup>™</sup> for Fibromyalgia: Preparing New Drug Application (NDA)

- Two Phase 3 trials completed with statistical significance on primary endpoint
- NDA filing expected 2H'24
- FDA decision on NDA approval expected 2H'25



### **Marketed Products**

 Zembrace® and Tosymra® indicated for the treatment of acute migraine



### **Pipeline**

- Phase 2 biologic cocaine antidote, FDA "Breakthrough Therapy Designation"
- Phase 1 anti-CD40L monoclonal antibody to prevent organ transplant rejection



## **Strategic Partnerships**

 With government institutions, world-class academic & research organizations



## **Internal Capabilities**

- Commercial prescription drug sales
- R&D and clinical-trial scale manufacturing



## **Key Clinical Programs**

Molecule*	Indication	Phase 1	Phase 2	Phase 3	NDA Submission
	Fibromyalgia	Statistically Significant Phase 3 Topline Results Reported 4Q'23		Submission expected 2H'24	
Tonmya™ TNX-102 SL	Long COVID		2 Topline Results		
Cyclobenzaprine HCl Protectic® Sublingual Tablets		Reported 3Q'23			La La Carte
Oubilityuai tablets	Acute Stress Disorder	Phase 2 Study** Start Expected 2Q'24			
TNX-1300		Phas	se 2 Study Start		A
Cocaine Esterase NIDA Funded	Cocaine Intoxication	Expected 2Q'24			
TNX-2900					1/2
Intranasal Potentiated Oxytocin FDA Orphan Drug and Rare Pediatric Disease Designation	han Drug and Rare Pediatric		2 Ready		
TNX-1500	Organ Transplant Rejection/	Phase 1 Stu			
Anti-CD40L mAb	Autoimmune Conditions	Ongoing expected 3Q'24		199	

<sup>\*</sup>All of Tonix's product candidates are investigational new drugs or biologics and none has been approved for any indication.

<sup>\*\*</sup>Investigator-initiated study



## **About Fibromyalgia**

Fibromyalgia is a <u>chronic pain disorder</u> resulting from amplified sensory and pain signaling within the CNS<sup>1</sup>

Fibromyalgia is a <u>syndrome</u> comprised of the <u>symptoms</u>: chronic widespread pain, <u>nonrestorative sleep</u>, and fatigue









Fibromyalgia is considered a chronic overlapping pain condition (COPC)

- the only COPC with any FDA-approved drugs<sup>3</sup>

Fibromyalgia is the prototypic nociplastic syndrome





## Fibromyalgia is a Large, Underserved and Dissatisfied population

- ~10 million U.S. adults are affected predominantly women<sup>1,2</sup>
  - Debilitating and life altering condition
  - Significant economic cost
- Patients are dissatisfied, despite three FDA approved drugs<sup>3,4</sup>
  - Average patient has 20 physician office visits per year<sup>2</sup>
  - Typical for patients to rotate between drugs<sup>3</sup>
  - Polypharmacy (multiple drugs at the same time) common<sup>3</sup>
  - Estimated that >22 million prescriptions are issued for the treatment of fibromyalgia (on- and off-label usage) each year<sup>5,6</sup>
- Prescription opiate use declining because of availability
  - Unknown number of patients using 'street drugs'
- No new Rx product since 2009



<sup>&</sup>lt;sup>1</sup>American College of Rheumatology (<u>www.ACRPatientInfo.org</u> accessed May 7, 2019) – prevalence rate of 2-4% for U.S. adult population (~250 million)

<sup>&</sup>lt;sup>2</sup>Vincent A, et al. *Arthritis Care Res (Hoboken)*. 2013 65(5):786-92. doi: 10.1002; diagnosed prevalence rate was 1.1% of adult population or 50% of the prevalent population

<sup>&</sup>lt;sup>3</sup>Robinson RL, et al. *Pain Med*. 2012 13(10):1366-76. doi: 10.1111; ; 85% received drug treatment

<sup>&</sup>lt;sup>4</sup>The three drugs with FDA approval for the treatment of fibromyalgia: Pregabalin (Lyrica); Duloxetine (Cymbalta); Milnacipran (Savella)

<sup>&</sup>lt;sup>5</sup>Product sales derived from IMS MIDAS; IMS NDTI used to factor usage for fibromyalgia; data accessed April 2015.

<sup>&</sup>lt;sup>6</sup>Market research by Frost & Sullivan, commissioned by Tonix, 2011



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



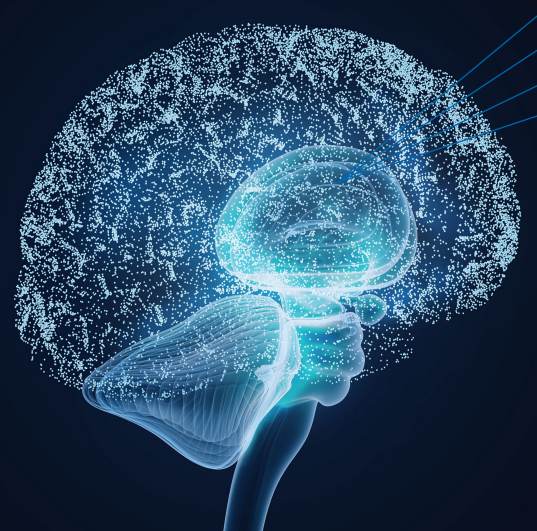
## Tonmya™ (TNX-102 SL, Cyclobenzaprine HCl Sublingual Tablets)¹

- Non-opioid analgesic designed for long-term daily bedtime use in fibromyalgia patients
  - Targets non-restorative sleep
  - No recognized risk for abuse
- Proprietary, sublingual formulation of cyclobenzaprine designed to optimize delivery and absorption
  - Issued patents expected to provide exclusivity to 2034/2035
  - Protectic® formulation based on eutectic composition of matter
- Improves <u>sleep quality</u>, does not increase <u>sleep quantity</u>:
  - Not a traditional hypnotic or sedative



# TNX-102 SL: Unique MOA Facilitates Restorative Sleep Centrally Acting Analgesic

Potent binding and antagonist activities at four key receptors facilitate *restorative sleep* 



- serotonergic-5-HT2A
- adrenergic-α1
- histaminergic-H1
- muscarinic-M1

## Key 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
- Not scheduled nor with recognized abuse potential

Issued patents expected to provide exclusivity to 2034/2035

Protectic® formulation based on eutectic composition of matter





## The Third Type of Pain: Nociplastic Pain<sup>1</sup>

#### **Nociplastic syndrome includes:**

- (1) widespread pain
- (2) fatigue
- (3) sleep disturbances
- (4) cognitive dysfunction ("brain fog")

## 

### **Nociplastic Pain**

Examples: I
Fibromyalgia /
ME/CFS per
Migraine
Irritable Bowel
Syndrome

Mechanism:
Altered pain
perception in the
brain

**Pathological Pain** 

### **Neuropathic Pain**

Examples: Sciatica Shingles Mechanism:
Impingement,
lesion or
inflammation of
nerve



# Fibromyalgia is Believed to Result from Chronic Pain or Prior Stress Experiences

#### The pain system evolved to detect acute pain

• The body's "check engine" light

# Chronic pain breaks down the system that determines whether a sensory experience is painful

- Chronic pain results in nociplastic syndromes
- Nociplastic syndrome was formerly known as "Central and Peripheral Sensitization"

## **Chronic Overlapping Pain Conditions** (COPCs) are Nociplastic Syndromes:

- Fibromyalgia
- ME/CFS
- Migraine
- Irritable Bowel Syndrome
- Endometriosis
- Low Back Pain

## Stresses that may precede or precipitate FM include:

#### Chronic nociceptive pain

• e.g., osteoarthritis

#### Chronic neuropathic pain

• *e.g.*, diabetic neuropathy

#### Infectious

• *e.g.*, viral illness

#### Cancer

• *e.g.*, breast cancer

#### Chemical

• e.g., cancer chemotherapy

#### **Traumatic**

• e.g., motor vehicle accident

#### Physiologic

• e.g., disturbed sleep





### Fibromyalgia is a common chronic disease<sup>1</sup>

Chronic pain syndrome that persists for years or decades

## No animal model is recognized for nociplastic syndromes or its component symptoms

- Widespread pain
- Fatigue
- Sleep disturbance
- Cognitive impairment

## Nociplastic symptoms are subjective

Humans need to report symptoms using scales

## Clinical trials measuring subjective symptoms are challenging

- Placebo response is typically observed
- Long-term therapy means requires long-term tolerability





## **Common Chronic Conditions are a Challenge for Society**

## The Opiate Crisis in the U.S. was driven by mistreatment of chronic pain, which was often nociplastic pain

- The epidemic of prescription pain killers was addressed by regulations which limited the availability of opiates
- Mandy individuals who are opiate dependent have transitioned to illegal street heroin and fentanyl
- Illegal drugs contribute to homelessness

## There is an unmet need for non-opiate analgesics that address nociplastic pain

No new drug for fibromyalgia has been approved since 2009







#### **Prevalence**

• One of the more common chronic pain disorders (2-4% of US Population)<sup>1</sup>

### **Diagnosed population**

- Large population but underdiagnosed<sup>2</sup> relative to prevalence rate
- Majority receive drug treatment<sup>3</sup>

#### **Treatment Pattern**

- Polypharmacy the norm average 2.6 drugs/patient<sup>3</sup>
- Rotation through therapy common: average ~5 drugs/year<sup>3</sup>
- Estimated that >22 million prescriptions are issued for the treatment of fibromyalgia (on- and off-label usage) each year<sup>4,5</sup>

#### **Unmet Need**

Majority of patients do not respond or cannot tolerate therapy<sup>6</sup>



<sup>&</sup>lt;sup>1</sup>American College of Rheumatology (<u>www.ACRPatientlnfo.org</u> accessed May 7, 2019) – prevalence rate of 2-4% for U.S. adult population (~250 million)

<sup>&</sup>lt;sup>2</sup>Vincent et al., 2013; diagnosed prevalence rate was 1.1% of adult population or 50% of the prevalent population

<sup>&</sup>lt;sup>3</sup>Robinson, et al., 2012; 85% received drug treatment

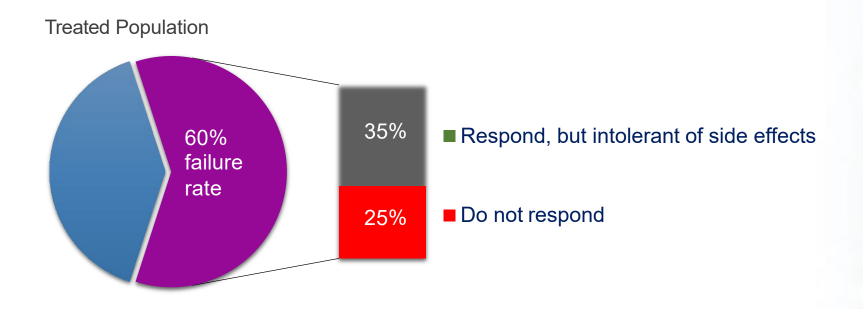
<sup>&</sup>lt;sup>4</sup>Vincent et al, Arthritis Care Res 2013;65:786

<sup>&</sup>lt;sup>5</sup>Product sales derived from IMS MIDAS; IMS NDTI used to factor usage for fibromyalgia; data accessed April 2015.

<sup>&</sup>lt;sup>6</sup>Market research by Frost & Sullivan, commissioned by Tonix, 2011

# Fewer than Half of Those Treated for Fibromyalgia Receive Sustained Benefit from the Three FDA-Approved Drugs<sup>1</sup>

- The treatment objective is to restore functionality and quality of life by broadly improving symptoms while avoiding significant side effects
- The majority fail therapy due to lack of a response or poor tolerability<sup>2</sup>





<sup>&</sup>lt;sup>1</sup> The three drugs with FDA approval for the treatment of fibromyalgia: Pregabalin (Lyrica); Duloxetine (Cymbalta); Milnacipran (Savella)

<sup>&</sup>lt;sup>2</sup> Market research by Frost & Sullivan, commissioned by Tonix (2011)



## Current FDA-Approved Fibromyalgia Drugs were Repurposed<sup>1</sup>

## Human investigation was required to find drugs that improve pain in fibromyalgia

• No current product addresses pain, poor sleep and fatigue

Drug		Lyrica® - Pfizer	Cymbalta® - Lilly Savella® - AbbVie
Initial Indication Sought		Epilepsy	Depression
Class		Gabapentinoid	SNRI
Mechanism		Slow neuron firing	Block NE reuptake
Fibromyalgia Activity	Pain	+	+
	Sleep	+	-
	Fatigue	-	+
Tolerability Issues	Sleep	-	+
	Fatigue	+	-
		Weight gain	Blood Pressure increases
			Sexual function impairment
			GI issues



# Large Need for New Fibromyalgia Therapies that Provide Broad Symptom Improvement with Better Tolerability

- Currently-approved medications may have side effects that limit long-term use<sup>1</sup>
  - Many patients skip doses or discontinue altogether within months of treatment initiation
- Medication-related side effects may be similar to fibromyalgia symptoms
- High rates of discontinuation, switching and augmentation
  - Attempt to treat multiple symptoms and/or avoid intolerable side effects
  - Average of 2-3 medications usedsimultaneously<sup>2</sup>
  - The typical patient has tried six different medications<sup>3</sup>
- Substantial off-label use of narcotic painkillers and prescription sleep aids<sup>3</sup>
  - Among those diagnosed, more than one-third have used prescription opioids as a means of treatment4
- Tonmya<sup>™</sup> (TNX-102 SL) is a non-opioid, centrally-acting analgesic that could provide a new therapeutic option for fibromyalgia patients

TONIX
PHARMACEUTICALS

<sup>&</sup>lt;sup>2</sup> Robinson RL et al, Pain Medicine 2012;13:1366.

<sup>&</sup>lt;sup>3</sup> Patient Trends: Fibromyalgia", Decision Resources,2011.

<sup>&</sup>lt;sup>4</sup> Berger A, Dukes E, Martin S, Edelsberg J, Oster G, Int J Clin Pract, 2007; 61(9):1498–1508.

## Fibromyalgia Program Status

Tonmya™\* (TNX-102 SL)

**Fibromyalgia** 

Statistically Significant 2<sup>nd</sup> Phase 3 Topline Results
Reported 4Q'23

Cyclobenzaprine Protectic® Sublingual Tablets



First pivotal Phase 3 study (RELIEF) reported – December 20201



Second Phase 3 study (RALLY) missed primary endpoint – July 2021



Confirmatory pivotal Phase 3 study (RESILIENT) reported – December 2023

## **Next Steps:**

- Type B Pre-NDA meeting scheduled with FDA in 2Q'24
- NDA filing expected 2H'24
- FDA decision on NDA approval expected 2H'25

<sup>\*</sup>Tonmya™ is conditionally accepted by the U.S. Food and Drug Administration (FDA) as the tradename for TNX-102 SL for the management of fibromyalgia. Tonmya has not been approved for any indication.



## Tonmya<sup>™</sup> (TNX-102 SL): Phase 3 *RESILIENT* Study Design



### **General study characteristics:**

- Randomized, double-blind, multicenter, placebo-controlled study in fibromyalgia
- 33 U.S. sites enrolled 457 participants with fibromyalgia as defined by 2016 Revisions to the 2010/2011 FM Diagnostic Criteria<sup>1</sup>

### **Primary Endpoint:**

Change from baseline to Week 14 (TNX-102 SL vs. placebo) in weekly averages of daily diary average pain severity score

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: NCT05273749

Study Title: A Phase 3 Study to Evaluate the Efficacy and Safety of TNX-102 SL

Taken Daily in Patients With Fibromyalgia (RESILIENT)

Trial ID: TNY-CY-F307 ('RESILIENT')

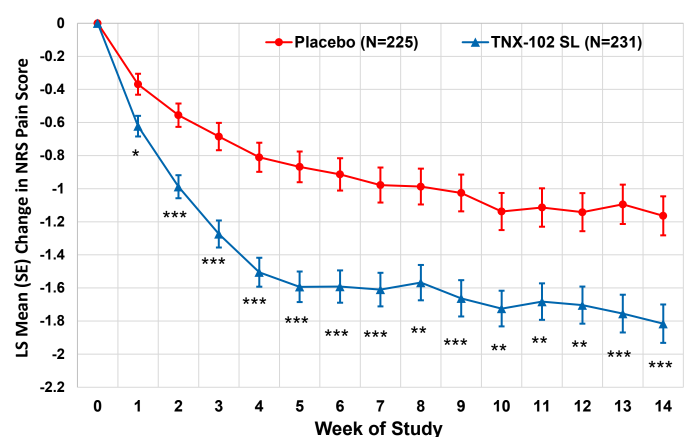


# RESILIENT Primary Outcome Measure Reduction in Widespread Pain





### Weekly Average of Daily Diary NRS Ratings of Average Pain Over Prior 24 Hours



\*p<0.01; \*\*p<0.001; \*\*\*p<0.0001

Week 14 LS mean (SE) change from baseline for TNX-102 SL -1.82 (0.12) and for placebo -1.16 (0.12); LSMD from placebo -0.65 (0.16); p=0.00005#

\*Based on Mixed Model Repeated Measures with Multiple Imputation, with fixed categorical effects of treatment, center, study week, and treatment by study week interaction, as well as baseline value and baseline value-by-study week interaction. Abbreviations: LS, least squares; LSMD, least squares mean difference; NRS, numerical rating scale; SE, standard error





## **Summary of Key Pre-Specified Secondary Outcome Measures**

Rating Scale	<u>Week 14</u>	<u>Met**</u>
Patient Global Impression of Change (PGIC)	<i>p</i> < 0.001	$\checkmark$
Fibromyalgia Impact Questionnaire - Symptoms	<i>p</i> < 0.001	<b>√</b>
Fibromyalgia Impact Questionnaire - Function	p = 0.001	<b>√</b>
PROMIS Sleep Disturbance	<i>p</i> < 0.001	$\checkmark$
PROMIS Fatigue	<i>p</i> < 0.001	$\checkmark$
Weekly average of daily Sleep Quality scores	p < 0.001	✓

<sup>\*</sup>In order of statistical serial gate-keeping hierarchy (or, "waterfall") to control overall Type 1 error \*\*Statistical significance met







## **RESILIENT Pre-Specified Primary Endpoint**

## **Summary**<sup>1</sup>

- Tonmya<sup>™</sup> (TNX-102 SL) demonstrated statistically significant improvement in mean weekly pain scores over placebo at Week 14
- <u>P-value of 0.00005</u> is *highly* statistically significant

## **Additional Findings**

- Effect size 0.38
- All pre-specified sensitivity analyses of the primary endpoint show statistical significance (p ≤ 0.001)
- Rapid onset of action: p-values <0.01 at each weekly time point, including Week 1</li>



# RESILIENT



## **RESILIENT** Safety Summary

Among participants randomized to Tonmya™ (TNX-102 SL) and to placebo, 81.0% and 79.6%, respectively, completed the study

Tonmya™ (TNX-102 SL) was generally well tolerated with an adverse event (AE) profile comparable to prior fibromyalgia studies

- No new safety signals were observed
- AE-related study discontinuations occurred in 6.1% and 3.6% of patients in the TNX-102 SL and placebo groups, respectively
- Events rated as mild or moderate made up 97.2% of AEs on placebo and 99.1% on TNX-102 SL
- As observed in prior studies with TNX-102 SL, oral administration site AEs were higher in TNX-102 SL than placebo, 42.9% and 10.2%, respectively
  - Most common oral AEs were oral hypoaesthesia, product taste abnormal, oral paraesthesia, and tongue discomfort (see table on next slide)
  - Nearly all of these common oral AEs were temporally related to dosing and lasted <60 minutes</li>
- Serious Adverse Events (SAEs)
  - Three placebo participants experienced an SAE:
    - 1. Pneumonia, 2. Muscular weakness, and 3. Hypertension/Angina/Coronary Artery Disease
  - Two TNX-102 SL participants experienced an SAE
    - 1. Renal carcinoma deemed not related to study drug
    - 2. Acute pancreatitis with onset 14 days after completion of treatment phase, deemed 'possibly related'\* to study drug
      - Outcome: 'Recovered/Resolved'
      - \*Note: participant was non-compliant with end of treatment study visits, and the last dose before onset of SAE was not known at the time that relationship with study drug was assessed by Investigator and Sponsor



## **RESILIENT** Safety Summary

## Treatment-Emergent Adverse Events (TEAEs) at Rate of ≥ 3% in Either Treatment Group

System Organ Class Preferred Term	TNX-102 SL N=231	Placebo N=226	Total* N=457
Systemic Adverse Events			
COVID-19	10 (4.3%)	7 (3.1%)	17 (3.7%)
Somnolence	7 (3.0%)	3 (1.3%)	10 (2.2%)
Headache	7 (3.0%)	4 (1.8%)	11 (2.4%)
Oral Cavity Adverse Events			
Hypoaesthesia oral	55 (23.8%)	1 (0.4%)	56 (12.3%)
Product taste abnormal	27 (11.7%)	2 (0.9%)	29 (6.3%)
Paraesthesia oral	16 (6.9%)	2 (0.9%)	18 (3.9%)
Tongue discomfort	16 (6.9%)	0 (0.0%)	16 (3.5%)

\*Safety Population

#### Changes in Sexual Functioning Questionnaire short form (CSFQ-14) was a safety measure in the study

- In females, CSFQ-14 total score improved (indicating better sexual functioning) to a greater extent in the TNX-102 SL group compared with placebo, p=0.010
- Orgasm/Completion and Desire/Frequency were improved
- Potential tolerability advantage over pharmacotherapeutics with potent serotonin reuptake inhibition





## Tonmya<sup>™</sup> Showed Broad-Spectrum Activity and was Well Tolerated

		Lyrica®	Cymbalta® Savella®	Tonmya™
Activity	Pain	YES	YES	YES
	Sleep	YES	-	YES
	Fatigue	-	YES	YES
Systemic Tolerability Issues	Insomnia	-	+	-
	Fatigue	+	-	-
	Weight	+	-	-
	Blood Pressure	-	+	-
	Sexual function	-	+	-
	GI issues	-	+	-

- Tonmya showed activity in all three measures of pain, sleep, and fatigue
- Tonmya is not associated with any of the commonly reported side effects



## Tonmya<sup>™</sup> (TNX-102 SL): Fibromyalgia Cyclobenzaprine Protectic<sup>®</sup> 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., the majority of whom are women<sup>1</sup>
- 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

#### **DEVELOPMENT PROGRAM**

Market Entry: Fibromyalgia

**Status:** Pivotal Phase 3 study RELIEF statistically significant, *p*-value = 0.01<sup>2</sup>

Second Phase 3 study RALLY missed primary endpoint

Confirmatory Phase 3 study RESILIENT statistically significant, *p*-value = 0.00005

**Next Steps:** Type B Pre-NDA meeting with FDA scheduled for 2Q'24

Additional Indications: Fibromyalgia-type Long COVID, Acute Stress Disorder (ASD), PTSD, Agitation in Alzheimer's, Alcohol Use Disorder

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





## Planning for Tonmya<sup>™</sup> Launch and Marketing

#### Several companies that successfully developed CNS drugs have launched them

• Big Pharma wants the commercial launch de-risked before acquisition (e.g., Nurtec®)

Company		Mkt Cap <sup>1</sup>	Product	Indication	FDA Approval	Exit
Axsome	AXSM	\$4.4 B	Auvelity®	Depression	8/2022	
Biohaven	BHVN	\$3.8 B	Nurtec®	Migraine	2/2020	Sold to PFE for \$12 B in Oct 2022
IntraCellular	ITCI	\$7.1 B	Caplyta®	Schizophrenia	12/2019	
Supernus	SUPN	\$1.5 B	Oxtella-XR®	Seizures	1/2019	
Neurocrine	NBIX	\$13.2 B	Ingrezza®	Tardive Dyskinesia	4/2017	
Acadia	ACAD	\$4.1 B	Nuplazid®	Parkinson's psychosis	4/2016	

### To prepare for the launch of Tonmya, Tonix acquired two marketed Rx drugs: Zembrace® and Tosymra®

Both are indicated for the acute treatment of migraine



## Additional Potential Indications for Tonmya™ (TNX-102 SL)



## Fibromyalgia-Type Long COVID

Status: Phase 2

Phase 2 study (PREVAIL) completed

Topline results reported 3Q 2023

**Next Steps:** Meet with FDA



#### **Acute Stress Reaction/ Acute Stress Disorder**

- Phase 2 ready investigator-initiated study
- Department of Defense funded
- UNC will perform study
- Received IND clearance from FDA

**Next Steps:** Expect to start Phase 2 in 2Q 2024



# Role of Infections in Triggering Fibromyalgia or Chronic fatigue (CFS)-Like Illnesses

- Symptoms of Long COVID, like multi-site pain, fatigue and insomnia, are the hallmarks of chronic pain syndromes like fibromyalgia and myalgic encephalomyelitis/chronic fatigue syndrome (ME/CFS).
- In August 2022, the HHS released the *National Research Action Plan on Long COVID*<sup>1</sup> which endorses the connection between Long COVID and chronic fatigue syndrome.

## Infection initiates an autoreactive process, which affects several functions, including brain and energy metabolism<sup>2-7</sup>

- Infections can trigger any of these conditions in approximately 10% of exposed individuals
- The initial location of the infection determines the subsequent pain syndrome
- Any type of infectious diarrhea will trigger irritable bowel syndrome (IBS) in 10% to 20% of those exposed



<sup>&</sup>lt;sup>2</sup>Blomberg J, et al. Front Immunol. 2018;9:229. Published 2018 Feb 15.



<sup>&</sup>lt;sup>3</sup>Warren JW, et al. Urology. 2008;71(6):1085-1090.

<sup>&</sup>lt;sup>4</sup>Buskila D, et al. Autoimmun Rev. 2008;8(1):41-43.

<sup>&</sup>lt;sup>5</sup>Hickie I, et al. BMJ. 2006;333(7568):575.

<sup>&</sup>lt;sup>6</sup>Parry SD, et al. Am J Gastroenterol. 2003;98(9):1970-1975.

<sup>&</sup>lt;sup>7</sup>Halvorson HA, et al. Am J Gastroenterol. 2006;101(8):1894-1942.



Long COVID is broadly defined as signs, symptoms, and conditions that continue or develop after acute COVID-19 infection<sup>1</sup>









## Many Long-COVID symptoms overlap with core symptoms of fibromyalgia

and are hallmarks of other chronic pain syndromes like myalgic encephalomyelitis/chronic fatigue syndrome (ME/CFS)

19%

Long COVID occurs in approximately 19% of recovered COVID-19 patients<sup>2</sup>

40%

As many as 40% of Long COVID patients experience multi-site pain<sup>3,4</sup>



## **TNX-102 SL for Fibromyalgia-Type** Long COVID: Phase 2 PREVAIL Study Design



# **PREVAIL** Study

### **Study characteristics:**

- Randomized, double-blind, placebo-controlled study of TNX-102 SL in fibromyalgia-type Long COVID
- U.S. sites only, *completed enrollment of 63 patients*

### **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

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

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

Placebo once-daily at bedtime

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 Seguelae of SARS-CoV-2 Infection (PREVAIL)"

14 weeks

Next Steps: End of Phase 2 Meeting with FDA





## TNX-102 SL: Phase 2 PREVAIL Topline Results<sup>1</sup>

Did not meet the primary endpoint of multi-site pain reduction at Week 14

However, findings fulfill the objectives of proof-of-concept study, supporting the decision to advance the program based on a proposed primary endpoint using the PROMIS Fatigue scale

- TNX-102 SL showed robust effect size in improving fatigue and consistent activity across secondary measures of sleep quality, cognitive function, disability and Patient Global Impression of Change (PGIC)
- Was generally well tolerated with an adverse event (AE) profile comparable to prior studies with TNX-102 SL:
  - AE-related discontinuations were similar in drug and placebo arms
  - No new safety signals were observed

Fatigue is the signature symptom of Long COVID and has been identified as the dominant symptom contributing to disability<sup>2</sup>

- We observed numerical improvement in the PROMIS fatigue score (in RELIEF p=0.007 MMRM and in RALLY p=0.007 MMRM) in both prior Phase 3 studies of TNX-102 SL in fibromyalgia,
- We believe the results of PREVAIL, together with extensive data from studies in other chronic conditions<sup>3-5</sup>, makes PROMIS Fatigue a solid candidate for the primary endpoint of future Long COVID registrational studies



<sup>&</sup>lt;sup>2</sup>Walker S, *et al. BMJ Open* 2023;13:e069217. doi:10.1136/bmjopen-2022-069217



## Acute Stress Reaction (ASR) / Acute Stress Disorder (ASD)

ASR/ASD are acute stress conditions resulting from trauma which can affect both civilian and military populations.

### Large unmet need:

- According to National Center for PTSD, about 60% of men and 50% of women in the US are exposed least one traumatic experience in their lives<sup>1</sup>
- In the US alone, one-third of emergency department visits (40-50 million patients per year) are for evaluation after trauma exposures<sup>2</sup>

#### **Current standard of care:**

 No medications are currently available at or near the point of care to treat patients suffering from acute traumatic events and support long-term health





## **TNX-102 SL for ASR/ASD: Program Status**

Status: Expect to start Phase 2 in 2Q 2024; received IND clearance from FDA

Phase 2 Trial Funded by DoD grant to University of North Carolina (UNC)

- UNC Institute for Trauma Recovery awarded a \$3M grant from the Department of Defense (DoD)
- OASIS trial will build upon infrastructure developed through the UNC-led, \$40M AURORA initiative
  - AURORA study is a major national research initiative to improve the understanding, prevention, and recovery of individuals who have experienced a traumatic event
  - Supported in part by funding from the National Institutes of Health (NIH) and the health care arm of Google's parent company
     Alphabet
- Opportunity to investigate the correlation between motor vehicle collisions and the emergence of ASD and PTSD
- Supported by multiple clinical trials:
  - Phase 2 trial in military-related PTSD (AtEase or NCT02277704)
  - Phase 3 trial in military-related PTSD (HONOR or NCT03062540)
  - Phase 3 trial in primarily civilian PTSD (RECOVERY or NCT03841773)
- In each of these studies, early and sustained improvements in sleep were associated with TNX-102 SL treatment by the PROMIS sleep disturbance (SD) scale and the Clinician Administered PTSD Scale (CAPS-5) "sleep disturbance" item.

Together these studies provide preliminary evidence that TNX-102 SL is well-tolerated and may promote recovery from PTSD via a pharmacodynamic facilitation of sleep-dependent emotional memory processing



# TNX-102 SL for ASR/ASD: Phase 2 OASIS Study Design

#### **General study characteristics:**

- Randomized, double-blind, placebo-controlled study in Acute Stress Reaction (ASR) / Acute Stress Disorder (ASD)
- The proposed Optimizing Acute Stress reaction Interventions with TNX-102 SL (OASIS) trial will examine the safety and efficacy of TNX-102 SL to reduce adverse posttraumatic neuropsychiatric sequelae among patients presenting to the emergency department after a motor vehicle collision (MVC)
- The trial will enroll approximately 180 individuals who acutely experienced trauma at study sites across the US
- Participants will be randomized in the emergency department to receive a two-week course of either TNX-102 SL or placebo
- Investigator-initiated IND

#### **Objective:**

- Investigate the potential of Tonix's TNX-102 SL (cyclobenzaprine HCl sublingual tablets) to reduce the frequency and severity of the adverse effects of traumatic exposure, including acute stress reaction (ASR), acute stress disorder (ASD), and posttraumatic stress disorder (PTSD).
- ASR refers to the body's immediate response to trauma, whereas ASD is the short-term effects of trauma (within 1 month), and PTSD is the long-term effects of trauma (beyond 1 month)

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

Placebo once-daily at bedtime

2 weeks

\*First dose of TNX-102 SL 5.6 mg versus placebo taken in the emergency department, and then daily at bedtime to finish 2 weeks of treatment

# A Phase 2 Study to Evaluate the Efficacy and Safety of TNX-102 SL Taken Daily in Patients With ASR/ ASD (OASIS)

- Primary outcome measure: Acute Stress Disorder Scale (ASDS) assessed at 7 and 21 days post MVC
- Posttraumatic stress symptom severity assessed at 6 and 12 weeks post MVC using the PTSD Checklist for DSM-5 (PCL-5)
- Standardized survey instruments of sleep disturbances, anxiety and depression symptoms, general physical and mental health, and clinical global improvement also employed
- Detailed and brief neurocognitive assessments are performed from baseline to 12 weeks after MVC at specific timepoints throughout study participation period

# Tonmya<sup>™</sup> (TNX-102 SL): Patents and Patent Applications

#### U.S. Composition:\*

- A 75:25 cyclobenzaprine HCl mannitol eutectic (dependent claims add a basifying agent).
  - 5 US Patents (Expire November 2034)
  - 1 Pending US Application (Would expire November 2034)
- A composition of a cyclobenzaprine HCl and a basifying agent suitable for sublingual absorption.
  - 1 Pending US Application (Would expire June 2033)

#### U.S. Methods of Use\* (Specific Indications):

- Fibromyalgia
  - Pain, Sleep Disturbance, Fatigue
    - 1 Pending US Application (Would expire December 2041)
  - Early Onset Response
    - 1 Pending US Provisional Application (Would expire December 2044)
  - Depressive Symptoms
    - 1 Pending US Application (Would expire March 2032)
- Sexual Dysfunction
  - 1 Pending US Application (Would expire October 2041)
- PASC
  - 1 Pending US Application (Would expire June 2043)
- PTSD
  - 1 US Patent (Expires November 2030)
- Agitation (Dementia)
  - 1 US Patent (Expires December 2038)
  - 1 Pending US Application (Would expire December 2038)
- Alcohol Use Disorder
  - 1 Pending US Application (Would expire November 2041)

#### Foreign Filings

- Corresponding foreign patents have been filed and some have issued:
  - Composition (25 patents, 3 allowed applications, 16 pending applications)
  - Methods of Use (9 patents, 54 pending applications)







# **Tonix Medicines: Commercial-Stage Specialty Pharma Subsidiary**

- Tonix Medicines is a wholly-owned subsidiary of Tonix (NASDAQ: TNXP)
  - Currently marketing two products indicated for the treatment of acute migraine: Zembrace<sup>®</sup> SymTouch <sup>®</sup> and Tosymra<sup>®</sup>
  - ~16 M in net sales<sup>1</sup>
  - Nascent commercial organization
- Tonix Medicines is led by James (Jim) Hunter
  - Veteran pharma executive with a track record for growing early businesses
  - Hunter previously founded Validus with Tonix CEO, Dr. Lederman
- Tonix Medicines is preparing to launch Tonmya<sup>™</sup> (TNX-102 SL) for fibromyalgia
  - Fibromyalgia care is relatively concentrated to specialized providers
  - We believe prescribing physicians can be targeted effectively by a specialty sales force
  - Evolving landscape in commercial markets favors distribution channels such as specialty pharmacies



# **Two Marketed Proprietary Migraine Drugs**

#### **Non-oral Formulations of Sumatriptan**

# Zembrace® SymTouch® (sumatriptan injection) 3 mg<sup>1</sup>



# Tosymra® (sumatriptan nasal spray) 10 mg²



- Each indicated for the treatment of acute migraine with or without aura in adults
- Sumatriptan remains the acute migraine 'gold standard' treatment for many patients and continues to represent the largest segment of the market in terms of unit sales<sup>3</sup>
- Each may provide migraine pain relief in as few as 10 minutes for some patients<sup>1,2,4,5</sup>
- Patents to 2036 (Zembrace) and 2031 (Tosymra)

# Acquired from Upsher-Smith Laboratories which has managed care contracts covering ~200 M lives

Contract includes a transition period during which Tonix expects to secure its own contracts

#### **Tonix Medicines Commercial Subsidiary**

- Complete commercialization capability
  - Manage supply chain and contract manufacturer
  - Distribution
  - Trade, Managed Care & Government contracting
- Team of professionals including Sales & Marketing personnel

<sup>1</sup>Zembrace SymTouch [package insert]. Maple Grove, MN: Upsher-Smith Laboratories, LLC: February 2021 - For more information, talk to your provider and read the <u>Patient Information</u> and <u>Instructions for Use</u>. – Important Safety Information is provided in the appendix <sup>2</sup>Tosymra [package insert]. Maple Grove, MN: Upsher-Smith Laboratories, LLC: Feb 2021. For more information, talk to your provider and read the <u>Patient Information</u> and <u>Instructions for Use</u>. – Important Safety Information is provided in the appendix <sup>3</sup>Upsher-Smith Laboratories, LLC; Data On File, 2023

<sup>4</sup>Mathew NT, et al. Dose ranging efficacy and safety of subcutaneous sumatriptan in the acute treatment of migraine. US Sumatriptan Research Group. Arch Neurol. 1992;49(12):1271-1276.

<sup>5</sup>Wendt J, et al. A randomized, double-blind, placebo-controlled trial of the efficacy and tolerability of a 4-mg dose of subcutaneous sumatriptan for the treatment of acute migraine attacks in adults. Clinical Therapeutics. 2006;28(4):517-526.

Zembrace SymTouch and Tosymra are registered trademarks of Tonix Medicines, Inc. Intravail is a registered trademark of Aegis Therapeutics, LLC, a wholly owned subsidiary of Neurelis, Inc.





# **Zembrace and Tosymra Bypass the GI Tract**

#### Bypassing gastrointestinal (GI) tract is potential advantage for treating acute migraine

- GI absorption may be inconsistent in migraineurs due to gastric stasis (also called "gastroparesis")<sup>1-4</sup>
- Nausea and vomiting are symptoms of migraine<sup>5</sup> which can complicate oral treatment

#### **Existing intranasal products**

- Imitrex® nasal spray (sumatriptan)
- Migranal® (dihydroergotamine) nasal spray developed by Novartis, sold by Bausch Health

#### New intranasal product bringing attention to non-oral route

• Pfizer's Zavzpret® (zavegepant), FDA approved in March, 2023<sup>1</sup> is the first intranasal gepant





# **Pipeline Development Strategy**

#### Focusing on government and academic collaborations

- Validates Tonix's scientific expertise and technology
- Reduces internal spend
- Increases number of trials
- Potentially speeds time to market
- Grants, contracts, cost-sharing or "in-kind" arrangements



# **External Partnerships**

#### Government partners providing direct funding, cost sharing or in-kind support include:

- National Institutes of Health (NIH)
- National Institute of Allergy and Infectious Disease (NIAID)
  - TNX-1800 selected for Project NextGen
- National Institute on Drug Abuse (NIDA)
  - TNX-1300 for cocaine intoxication; Phase 2 study funding
- Department of Defense (DoD)
  - TNX-102 SL for ASD; Phase 2 study funding

#### Academic partners sponsoring clinical trials of Tonix's investigational drug products include:

- Massachusetts General Hospital (MGH)
- University of Washington
- University of North Carolina



# **Key Partnerships**

**TNX-1500:** ALLOGRAFT REJECTION





TNX-102 SL: ACUTE STRESS DISORDER





THE UNIVERSITY
of NORTH CAROLINA
at CHAPEL HILL

**TNX-1300: COCAINE INTOXICATION** 





TNX-1800: COVID-19 VACCINE





TNX-2900: PRADER-WILLI SYNDROME







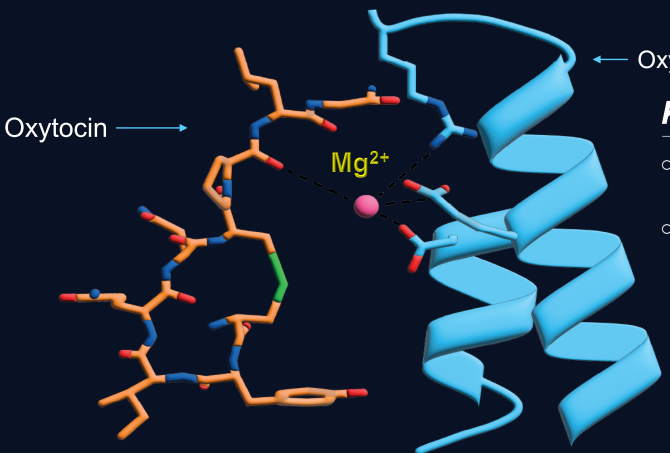




A novel, non-CGRP antagonist approach to treatment

# TNX-2900: Novel Formulation of Intranasal Oxytocin (OT) Potentiated with Magnesium

Magnesium is known to **potentiate the binding of OT** to its receptor<sup>1,2</sup>



Oxytocin receptor

# Key Differentiators

- Targeted intranasal delivery
  - Low systemic exposure
- When delivered via the nasal route, concentrates in trigeminal system
  - Binding of OT to receptors on neurons in trigeminal system inhibits release of CGRP and transmission of pain signals
  - Blocking CGRP release is a distinct mechanism compared with CGRP antagonist and anti-CGRP antibody drugs, which block the binding of CGRP to its receptor





# **About Prader-Willi Syndrome**

Prader-Willi Syndrome (PWS) is the most common genetic cause of life-threatening childhood obesity. PWS causes unhealthy behaviors around food<sup>1-4</sup>, consequences such as **obesity**, **type 2** diabetes, and cardiovascular disease<sup>1-5</sup>, and creates significant caretaker burden<sup>1-4</sup>

Rare genetic disease that afflicts 10-20 thousand individuals in the US

#### **Current standard of care:**

Human growth hormone treatment is FDA-approved for growth failure in PWS children

#### Large unmet need:

- Currently no cure, and no treatment for PWS-related hyperphagia
- Consequences can be life threatening obesity and cardiovascular disease are leading cause of death

\*TNX-2900 has been granted FDA Orphan Drug and Rare Pediatric Disease Designation, and received IND clearance by FDA for Phase 2 Trial

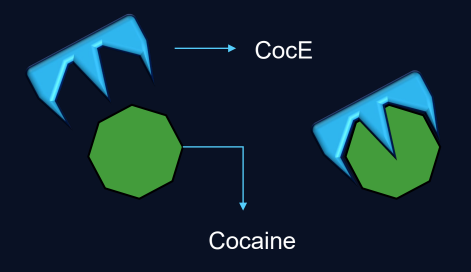


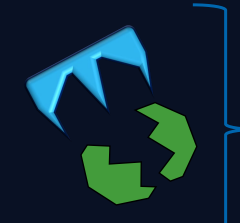


Fast acting antidote for life threatening cocaine intoxication

# TNX-1300: Recombinant Protein Rapidly Degrades Cocaine in the Bloodstream

Drops plasma exposure by 90% in 2 minutes





FDA Breakthrough Therapy Designation

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

# **Key Differentiators**

- Rapidly metabolizes cocaine within matter of minutes
- No other product currently on the market for this indication





### **About Cocaine Intoxication**

Over 5 million Americans reported current cocaine use in 2020, which is almost 2% of the population<sup>1</sup>. In 2021, more than 24,900 individuals in the US died from drug overdose deaths involving cocaine<sup>2</sup>

500k Over 500,000 emergency department visits for cocaine, annually<sup>3,4</sup>

#### **Current standard of care:**

 Patients are currently managed only by supportive care for the adverse effects of cocaine intoxication on the cardiovascular and central nervous systems

#### Large unmet need:

- No other product currently on the market for this indication
- TNX-1300 could significantly reduce the time and resources required for other detox services
- Potentially reduces the risk of morbidity and mortality



<sup>&</sup>lt;sup>1</sup>Substance Abuse and Mental Health Services Administration. (2021). Results from the 2020 National Survey on Drug Use and Health: Detailed Tables: Prevalence Estimates, Standard Errors, and Sample Sizes.

<sup>&</sup>lt;sup>2</sup> Centers for Disease Control and Prevention (CDC) - https://www.cdc.gov/nchs/nvss/vsrr/drugoverdose-data.htm

<sup>&</sup>lt;sup>3</sup>Substance Mental Health Services Administration, Drug Abuse Warning Network, 2011: National Estimates of Drug-Related Emergency Department Visits. HHS Publication No. (SMA) 13-4760, DAWN Series D-39. Rockville, MD: Substance Abuse and Mental Health Services Administration, 2013.

<sup>&</sup>lt;sup>4</sup> Drug Abuse Warning Network, 2011: Selected Tables of National Estimates of Drug-Related Emergency Department Visits. Rockville, MD: Center for Behavioral Health Statistics and Quality, SAMHSA, 2013.





## TNX-1500: Next Generation anti-CD40L mAb

Re-engineered to better modulate the binding of Fc \gamma\R and mitigate risk of thrombosis Clinical Stage of Phase 1 study completed – Topline expected 3Q 2024

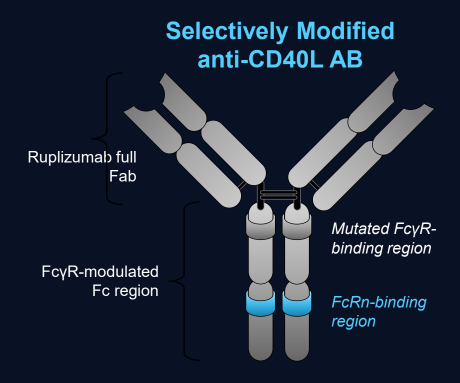
# **Key Differentiators**

Expected to deliver efficacy without compromising safety

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

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

Third Generation (TNX-1500): Re-engineered to better modulate the binding of FcγR.



Contains the full ruplizumab Fab and the engineered Fc region that modulates FcγR-binding, while preserving FcRn function





# **TNX-1500 Strategy and Status**

Proposed Initial Indication: Prevention of Allograft Rejection

Status: Phase 1 enrollment complete; data readout expected 3Q'24

- Collaborations ongoing with Mass General Hospital on heart and kidney transplantation in non-human primates
- Collaboration with Boston Children's on bone marrow transplantation in non-human primates

Next Steps: Initiate Phase 2 study in Kidney Transplant Recipients

- 2 Second Indication: Hematopoietic Cell Transplant (Bone Marrow Transplant)
  - Potential to reduce GvHD
- Third Indication (and beyond): Autoimmune Diseases (e.g., Multiple Sclerosis, Sjögen's Syndrome, Systemic Lupus Erythematosus)
  - These indications require large studies, but represent large target markets

Currently exploring strategic partnerships and out-licensing opportunities





### **TNX-1500 Preclinical Data and Publications**

#### Non-human Primate Kidney Allo-Transplantation

- TNX-1500 monotherapy consistently prevents kidney transplant rejection with no thrombosis observed
- April 2023 Publication: Lassiter, G., et al. (2023). TNX-1500, an Fc-modified Anti-CD154 Antibody, Prolongs Nonhuman Primate Renal Allograft Survival. *American Journal of Transplantation*. <a href="https://www.sciencedirect.com/science/article/pii/S1600613523003714">www.sciencedirect.com/science/article/pii/S1600613523003714</a>

#### **Non-human Primate Heart Heterotopic Allo-Transplantation**

- TNX-1500 monotherapy consistently prevents heart transplant rejection. Similar activity to chimeric hu5c8<sup>2</sup> during treatment phase in prior studies
- April 2023 Publication: Miura, S., et al. (2023) TNX-1500, an Fc-modified Anti-CD154 Antibody, Prolongs Nonhuman Primate
   Cardiac Allograft Survival. American Journal of Transplantation. <a href="https://www.sciencedirect.com/science/article/pii/S1600613523003969">www.sciencedirect.com/science/article/pii/S1600613523003969</a>

#### Non-Human Primate Kidney Xenograft Transplantation

- TNX-1500 therapy is part of a regiment to prevent rejection in kidney xenograft transplants
  - Anand, R.P., Layer, J.V., Heja, D. et al. (2023). Design and testing of a humanized porcine donor for xenotransplantation. *Nature*. <a href="https://www.nature.com/articles/s41586-023-06594-4">https://www.nature.com/articles/s41586-023-06594-4</a>
  - Kozlov, M. (2023). Monkey survives two years after gene-edited pig-kidney transplant. *Nature*.
     <a href="https://www.nature.com/articles/d41586-023-03176-2">https://www.nature.com/articles/d41586-023-03176-2</a>
  - Mohiuddin, M. (2023). Pig-to-primate organ transplants require genetic modifications of donor. *Nature*.
     <a href="https://www.nature.com/articles/d41586-023-02817-w">https://www.nature.com/articles/d41586-023-02817-w</a>



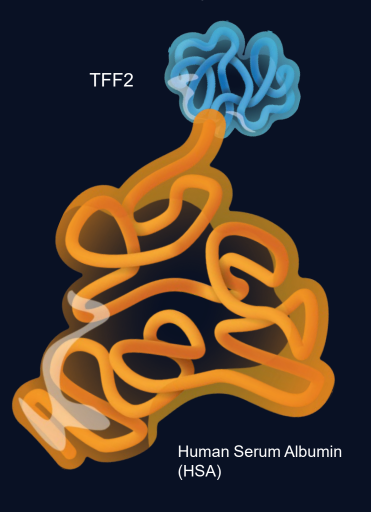
# TNX-1700

Recombinant Trefoil Factor Family Member 2 (rTFF2-HSA) Fusion Protein

Targeting the toxic tumor micro-environment

# **TNX-1700: Fighting Cancer by Targeting the Tumor Micro-Environment**

Suppresses myeloid-derived suppressor cells (MDSCs) and activates anti-cancer CD8+ T cells



## **Key Differentiators**

- Different MOA than checkpoint inhibitors
- o Potential synergy with anti-PD-1 or anti-PD-L1 monoclonal antibodies

#### **Preclinical Evidence**

- o mTNX-1700 (mTFF2-MSA fusion protein) and anti-PD-1 monotherapy each was able to evoke anti-tumor immunity in the MC38 model of colorectal cancer<sup>1</sup>
- mTNX-1700 augmented the anti-tumor efficacy of anti-PD-1 therapy in both the MC38 and the CT26.wt models<sup>1</sup>





## **About Gastric and Colorectal Cancer**

Gastric and colorectal cancer are both leading cancers in the US. Colorectal cancer is the 3rd leading cause of cancer-related deaths in both men and women.1

>1.3M

People living with colorectal cancer in the US<sup>2</sup>

>125k

People living with gastric cancer in the US<sup>3</sup>

#### **Current standard of care:**

- PD-1 blockade
  - However, gastric and colorectal cancer are relatively unresponsive

#### Large unmet need:

- Gastric and colorectal cancer have a relative 5-year survival rate of 35.7% and 65%, respectively
  - Despite advances in the field, patients are still in need of life saving treatment





# **Internal Development & Manufacturing Capabilities**



#### R&D Center (RDC): Frederick, MD

- Research advancing CNS and immunology drugs
- Accelerated development of vaccines and antiviral drugs against infectious diseases
- ~48,000 square feet, BSL-2 with some areas designated BSL-3



## Advanced Development Center (ADC): North Dartmouth, MA

- Development and clinical scale manufacturing of biologics
- ~45,000 square feet, BSL-2





# **Broad-Spectrum Antiviral Discovery Programs**

#### Host-directed antiviral discovery programs

#### **CD45** targeted therapeutics

- Small molecule therapeutics that reduce endogenous levels of CD45, a protein tyrosine phosphatase
- Reduction in CD45 protects against many viruses including the Ebola virus

#### **Cathepsin inhibitors**

- Small molecule therapeutics that inhibit **essential cathepsins** which are required by viruses such as coronaviruses and filoviruses to infect cells
- Activity as monotherapy and in combination with other antivirals

## Virus-directed antivirals discovery program

#### Viral glycan-targeted engineered biologics

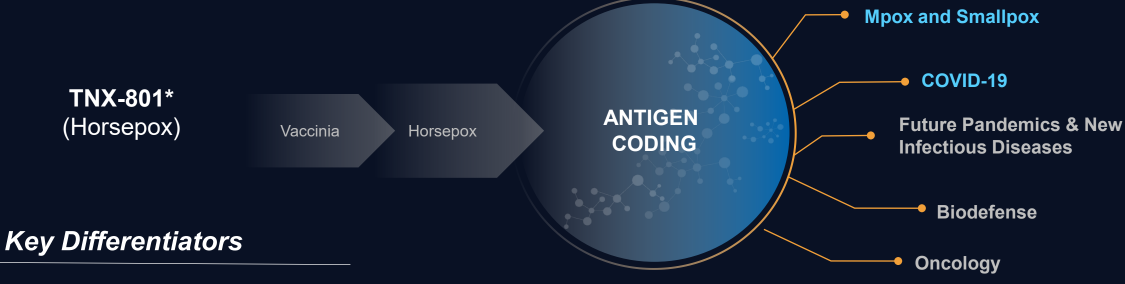
- Bind to viral densely branched high-mannose (DBH) glycans
- Neutralize circulating virus and stop the entry of the progeny virus into cells
- Antiviral activity against a broad range of RNA viruses
- Activity as monotherapy and in combination with other antivirals





# TNX-801: Recombinant Pox Vaccine (RPV) Platform Using Live Virus Technology

Cloned version of horsepox<sup>1</sup> purified from cell culture



#### Live virus vaccines are the most established vaccine technology

- Prevents forward transmission
- o Effective in eliciting durable or long-term immunity

#### **Economical to manufacture at scale**

- Low dose because replication amplifies dose in vivo
- Single administration

#### Standard refrigeration for shipping and storage





# TNX-1800: Designed to Express the SARs-CoV-2 Spike Protein

TNX-1800 (recombinant horsepox virus) is a live virus vaccine based on Tonix's TNX-801 that is designed to express the spike protein of the SARS-CoV-2 virus and to elicit a predominant T cell response

- Immunogenic and well tolerated<sup>1</sup>
- Showed promise in protecting animals from challenge with SARS-CoV-2 delivered directly into the lungs<sup>1</sup>

# Status: National Institute of Allergy and Infectious Diseases (NIAID) will conduct a Phase 1 clinical trial with TNX-1800

- First vaccine candidate using Tonix's live virus recombinant pox virus (RPV) platform technology to enter clinical trials
- "Project NextGen" is an initiative by the U.S. Department of Health and Human Services (HHS) to advance a
  pipeline of new, innovative vaccines and therapeutics for COVID-19. NIAID will be conducting clinical trials to
  evaluate several early-stage vaccine candidates, including TNX-1800
- Phase 1 study is designed to assess safety and immunogenicity in approximately 60 healthy adult volunteers
- Upon completion of the trial, NIAID and Tonix will assess the results and determine the next steps for the development of TNX-1800





# **Management Team**



**Seth Lederman, MD**Co-Founder, CEO & Chairman









**Gregory Sullivan, MD**Chief Medical Officer



New York State Psychiatric Institute



**Bradley Saenger, CPA**Chief Financial Officer











Jessica Morris
Chief Operating Officer









# Milestones: Recently Completed and Upcoming

#### **Financial Milestones**

4<sup>th</sup> Quarter 2023 Financing: \$144 M facility: \$30 M upfront

#### **Tonmya™ Milestones**

**1** 4<sup>th</sup> Quarter 2023 Statistically significant topline results of Phase 3 RESILIENT study for Tonmya<sup>™</sup> for the management of fibromyalgia

□ 2<sup>nd</sup> Quarter 2024 Type B Pre-NDA meeting with FDA for Tonmya<sup>™</sup> for fibromyalgia scheduled

□ 2<sup>nd</sup> Half 2024 Submit NDA to FDA for Tonmya<sup>™</sup> for fibromyalgia

#### **Other Key Program Milestones**

□ 2<sup>nd</sup> Quarter 2024 Initiate Phase 2 study of TNX-102 SL for acute stress disorder

□ 2<sup>nd</sup> Quarter 2024 Initiate Phase 2 study of TNX-1300 for the treatment of cocaine intoxication

□ 3<sup>rd</sup> Quarter 2024 Results of Phase 1 study of TNX-1500





# **Zembrace® Important Safety Information (1 of 2)**

Zembrace SymTouch (Zembrace) can cause serious side effects, including heart attack and other heart problems, which may lead to death. Stop use and get emergency help if you have any signs of a heart attack:

• Discomfort in the center of your chest that lasts for more than a few minutes or goes away and comes back; severe tightness, pain, pressure, or heaviness in your chest, throat, neck, or jaw; pain or discomfort in your arms, back, neck, jaw or stomach; shortness of breath with or without chest discomfort; breaking out in a cold sweat; nausea or vomiting; feeling lightheaded

Zembrace is not for people with risk factors for heart disease (high blood pressure or cholesterol, smoking, overweight, diabetes, family history of heart disease) unless a heart exam shows no problem.

Do not use Zembrace if you have:

- History of heart problems; narrowing of blood vessels to your legs, arms, stomach, or kidney (peripheral vascular disease); uncontrolled high blood pressure; hemiplegic or basilar migraines. If you are not sure if you have these, ask your provider.
- Had a stroke, transient ischemic attacks (TIAs), or problems with blood circulation; severe liver problems; taken any of the following medicines in the last 24 hours: almotriptan, eletriptan, frovatriptan, naratriptan, rizatriptan, ergotamines, dihydroergotamine; are taking certain antidepressants, known as monoamine oxidase (MAO)-A inhibitors or it has been 2 weeks or less since you stopped taking a MAO-A inhibitor. Ask your provider for a list of these medicines if you are not sure.
- An allergy to sumatriptan or any of the components of Zembrace

Tell your provider about all of your medical conditions and medicines you take, including vitamins and supplements.

Zembrace can cause dizziness, weakness, or drowsiness. If so, do not drive a car, use machinery, or do anything where you need to be alert.





# **Zembrace® Important Safety Information (2 of 2)**

#### Zembrace may cause serious side effects including:

- Changes in color or sensation in your fingers and toes; sudden or severe stomach pain, stomach pain after meals, weight loss, nausea or vomiting, constipation or diarrhea, bloody diarrhea, fever; cramping and pain in your legs or hips; feeling of heaviness or tightness in your leg muscles; burning or aching pain in your feet or toes while resting; numbness, tingling, or weakness in your legs; cold feeling or color changes in one or both legs or feet; increased blood pressure including a sudden severe increase even if you have no history of high blood pressure; medication overuse headaches from using migraine medicine for 10 or more days each month. If your headaches get worse, call your provider.
- Serotonin syndrome, a rare but serious problem that can happen in people using Zembrace, especially when used with anti-depressant medicines called SSRIs or SNRIs. Call your provider right away if you have: mental changes such as seeing things that are not there (hallucinations), agitation, or coma; fast heartbeat; changes in blood pressure; high body temperature; tight muscles; or trouble walking.
- Hives (itchy bumps); swelling of your tongue, mouth, or throat
- Seizures even in people who have never had seizures before

The most common side effects of Zembrace include: pain and redness at injection site; tingling or numbness in your fingers or toes; dizziness; warm, hot, burning feeling to your face (flushing); discomfort or stiffness in your neck; feeling weak, drowsy, or tired.

Tell your provider if you have any side effect that bothers you or does not go away. These are not all the possible side effects of Zembrace. For more information, ask your provider.

This is the most important information to know about Zembrace but is not comprehensive. For more information, talk to your provider and read the <u>Patient Information</u> and <u>Instructions for Use</u>. You can also visit <u>www.upsher-smith.com</u> or call 1-888-650-3789. For full Prescribing Information, visit: <a href="https://dailymed.nlm.nih.gov/dailymed/drugInfo.cfm?setid=6e5b104f-2b9e-416e-92fb-ef1bdaea867d">https://dailymed.nlm.nih.gov/dailymed/drugInfo.cfm?setid=6e5b104f-2b9e-416e-92fb-ef1bdaea867d</a>

You are encouraged to report adverse effects of prescription drugs to the FDA. Visit www.fda.gov/medwatch or call 1-800-FDA-1088.

Zembrace is a prescription medicine used to treat acute migraine headaches with or without aura in adults who have been diagnosed with migraine.

Zembrace is not used to prevent migraines. It is not known if it is safe and effective in children under 18 years of age.



# **Tosymra® Important Safety Information (1 of 2)**

Tosymra® can cause serious side effects, including heart attack and other heart problems, which may lead to death. Stop Tosymra and get emergency medical help if you have any signs of heart attack:

Discomfort in the center of your chest that lasts for more than a few minutes or goes away and comes back; severe
tightness, pain, pressure, or heaviness in your chest, throat, neck, or jaw; pain or discomfort in your arms, back, neck, jaw,
or stomach; shortness of breath with or without chest discomfort; breaking out in a cold sweat; nausea or vomiting; feeling
lightheaded

Tosymra is not for people with risk factors for heart disease (high blood pressure or cholesterol, smoking, overweight, diabetes, family history of heart disease) unless a heart exam is done and shows no problem.

Do not use Tosymra if you have:

- History of heart problems; narrowing of blood vessels to your legs, arms, stomach, or kidney (peripheral vascular disease); uncontrolled high blood pressure; severe liver problems; hemiplegic or basilar migraines. If you are not sure if you have these, ask your healthcare provider.
- Had a stroke, transient ischemic attacks (TIAs), or problems with blood circulation; taken any of the following medicines in the last 24 hours: almotriptan, eletriptan, frovatriptan, naratriptan, rizatriptan, ergotamines, or dihydroergotamine. Ask your provider if you are not sure if your medicine is listed above
- are taking certain antidepressants, known as monoamine oxidase (MAO)-A inhibitors or it has been 2 weeks or less since you stopped taking a MAO-A inhibitor. Ask your provider for a list of these medicines if you are not sure
- An allergy to sumatriptan or any ingredient in Tosymra

Tell your provider about all of your medical conditions and medicines you take, including vitamins and supplements. Tosymra can cause dizziness, weakness, or drowsiness. If so, do not drive a car, use machinery, or do anything where you need to be alert.





# **Tosymra® Important Safety Information (2 of 2)**

#### Tosymra may cause serious side effects including:

- Changes in color or sensation in your fingers and toes; sudden or severe stomach pain, stomach pain after meals, weight loss, nausea or vomiting, constipation or diarrhea, bloody diarrhea, fever; cramping and pain in your legs or hips, feeling of heaviness or tightness in your leg muscles, burning or aching pain in your feet or toes while resting, numbness, tingling, or weakness in your legs, cold feeling or color changes in one or both legs or feet; increased blood pressure including a sudden severe increase even if you have no history of high blood pressure; medication overuse headaches from using migraine medicine for 10 or more days each month. If your headaches get worse, call your provider.
- Serotonin syndrome, a rare but serious problem that can happen in people using Tosymra, especially when used with anti-depressant medicines called SSRIs or SNRIs. Call your provider right away if you have: mental changes such as seeing things that are not there (hallucinations), agitation, or coma; fast heartbeat; changes in blood pressure; high body temperature; tight muscles; or trouble walking.
- Seizures even in people who have never had seizures before

The most common side effects of Tosymra include: tingling, dizziness, feeling warm or hot, burning feeling, feeling of heaviness, feeling of pressure, flushing, feeling of tightness, numbness, application site (nasal) reactions, abnormal taste, and throat irritation.

Tell your provider if you have any side effect that bothers you or does not go away. These are not all the possible side effects of Tosymra. For more information, ask your provider.

This is the most important information to know about Tosymra but is not comprehensive. For more information, talk to your provider and read the Patient Information and Instructions for Use. You can also visit www.upsher-smith.com or call 1-888-650-3789. For full Prescribing Information, visit: https://dailymed.nlm.nih.gov/dailymed/drugInfo.cfm?setid=015a5cf9-f246-48bc-b91e-cd730a53d8aa

You are encouraged to report negative side effects of prescription drugs to the FDA. Visit www.fda.gov/medwatch, or call 1-800-FDA-1088.

Tosymra is a prescription medicine used to treat acute migraine headaches with or without aura in adults.

Tosymra is not used to treat other types of headaches such as hemiplegic or basilar migraines or cluster headaches.

Tosymra is not used to prevent migraines. It is not known if Tosymra is safe and effective in children under 18 years of age.

