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): March 5, 2019

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.)

509 Madison Avenue, Suite 306, New York, New York 10022 (Address of principal executive offices) (Zip Code)

Registrant's telephone number, including area code: (212) 980-9155

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))

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. \Box

Item 7.01 Regulation FD Disclosure.

Tonix Pharmaceuticals Holding Corp. the "Company") updated its investor presentations, which are 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. Copies of the investor presentations are filed as Exhibits 99.01, 99.02 and 99.03, and incorporated by reference in, this report.

Item 9.01 Financial Statements and Exhibits.

(d)	Exhibit No.	Description	
	99.01 99.02 99.03	<u>Corporate Presentation by the Company for March 5, 2019 (Long Form)</u> <u>Corporate Presentation by the Company for March 5, 2019 (Short Form)</u> <u>Corporate Presentation by the Company for March 5, 2019 (Abbreviated Form)</u>	

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.

Date: March 5, 2019

By: <u>/s/ Seth Lederman</u> Seth Lederman Chief Executive Officer

Exhibit 99.01





March 2019

Version P0161 3-5-19 (Doc 0439)



Cautionary Note on Forward-Looking Statements

2

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, substantial competition; 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 risks related to failure to obtain U.S. Food and Drug Administration clearances or approvals and noncompliance with its regulations. 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, 2017, as filed with the Securities and Exchange Commission (the "SEC") on March 9, 2018, and periodic 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.



Tonix Pharmaceuticals

Who we are:

 A clinical stage pharmaceutical company dedicated to developing innovative treatments for patients and making meaningful contributions to society 3

What we do:

- · Target therapeutics with high need for improvement
 - Conditions with no or ineffective treatments
 - Significant patient segments not well served by existing therapies
- Develop innovative treatment options with possibility to be a "game changer"
 - Scientifically unique and innovative
 - Supported by strong scientific rationale
 - Confirmed by clinical evidence and published literature
 - Utilize proven regulatory pathway and established clinical endpoint
 - Built on a foundation of proprietary intellectual property



ų	Posttraumatic Stress Disorder (PTSD) – Lead program; new bedtime
02 SL zaprir gual ets	 treatment - Tonmya®1 P302/RECOVERY Phase 3 clinical study with Week 4 primary endpoint to initiate in 1Q2019 Results from 2 efficacy studies improve the new Phase 3 study design New Phase
NX-1(loben sublin Tabl	P302/RECOVERY study design features accepted by the FDA ² Agitation in Alzheimer's disease (AAD)
Cycl	• IND ³ ready to support Phase 2 potential pivotal efficacy study Fibromyalgia Syndrome (FM)
	 IND³ ready to support Phase 3 potential pivotal efficacy study
Pipeline	TNX-601 ⁴ – Daytime PTSD treatment and treatment of neurocognitive dysfunction from corticosteroids • Pre-IND candidate; nonclinical development ongoing
Pi	TNX-801 ⁵ - Smallpox-preventing vaccine candidate Efficacy demonstrated in mouse model

is ¹ Tommya has been conditionally accepted by the U.S. FDA as the proposed an investigational new drug and has not been approved for any indication. ² FDA Meeting Minutes (November 26, 2018) ³ IND- Investigational New Drug Application ⁴ Tianeptine explate ⁵ Synthesized live horsepox virus ⁽⁸⁾ 2 ingu



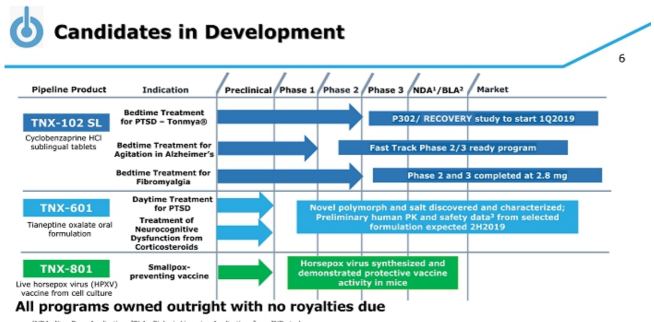
5

Sleep disturbances are associated with a constellation of disorders

- · Considered co-morbid or a key symptom in these disorders
- · Believed to have a role in the onset, progression and severity of these disorders

The focus of TNX-102 SL development is both unique and innovative

- Testing the therapeutic benefit of sleep ('sleep quality')
 - Restorative sleep...in contrast to time spent sleeping ('sleep quantity')
- Targeting clinical conditions for which improved sleep quality may have a therapeutic benefit
 - Reduction in disease-specific symptoms, with sleep improvement as a secondary endpoint



¹NDA- New Drug Application; ²BLA -Biologic Licensing Application; ³non-IND study (© 2019 Tonix Pharmaceuticals Holding Corp.

Tonmya: a Potential Bedtime Treatment for PTSD

First investigational new drug to show treatment effect in military-related PTSD in two potential pivotal efficacy studies

Phase 2 study (P201/AtEase) showed Tonmya 5.6 mg had a strong signal of treatment effect at Week 12 as measured by CAPS-51

7

- Phase 3 study (P301/HONOR) provided evidence of effectiveness as early as 4 weeks after treatment but diminished over time due to high placebo response
 - Retrospective analysis showed persistent effectiveness at Week 12 in subgroup with trauma ≤9 years from screening
- · Both studies can be used as supportive evidence of efficacy and safety for Tonmya NDA submission
- No serious or unexpected adverse events related to Tonmya were reported

FDA feedback and acceptance on new Phase 3 study (P302/RECOVERY) received in November² Patent protection through 2034 in U.S.³

Composition of matter patent for transmucosal delivery of cyclobenzaprine

Novel mechanism targets sleep quality

Memory processing during sleep is important to recovery from PTSD

CAPS-5 = Clinician-Administered PTSD Scale for DSM-5
 FDA Meeting Minutes, November 26, 2018; ³U.S. Patent No. 9,636,408 for eutectic proprietary ProtecticTM formulation





Active ingredient is cyclobenzaprine, which is structurally related to tricyclic antidepressants

- Cyclobenzaprine interacts with receptors that regulate sleep quality: 5-HT_{2A}, α_1 -adrenergic and histamine H₁ receptors
- Cyclobenzaprine does <u>NOT</u> interact with the same receptors as traditional hypnotic sleep drugs, benzodiazepines or nonbenzodiazepines that are associated with retrograde amnesia
- Cyclobenzaprine-containing product was approved 40 years ago and current labeling (May 2016) indicates no abuse or dependence concern

Tonmya NDA can be filed without drug abuse and dependency assessment studies

· Discussed at March 9, 2017 meeting with the FDA



TNX-102 SL Intellectual Property – U.S. Protection until 2034



- United States Patent and Trademark Office (USPTO) issued U.S. Patent No. 9,636,408 in May 201 7U.S. Patent No. 9,956,188 in May 2018 and U.S. Patent No. 10,117,936 in November 2018
- Japanese Patent Office (JPO) issued Japanese Patent No. 6310542 in March 2018
- New Zealand Intellectual Property Office (NZIPO) issued New Zealand Patent No. 631152 in May 2017

9

37 patent applications pending (2 allowed (US and South Africa))

Pharmacokinetics (PK) : Protection expected to 2033

- JPO issued Japanese Patent No. 6259452 in December 2017
- NZIPO issued New Zealand Patent No. 631144 in March 2017
- Taiwanese Intellectual Property Office issued Taiwanese Patent No. I590820 in July 2017
- · 21 patent applications pending (1 allowed (Australia))

Method of use for active ingredient cyclobenzaprine : Protection expected to 2030

- European Patent Office issued European Patent No. 2 501 234B1 in September 2017 (validated in 38 countries). Opposition filed in June 2018
- USPTO issued U.S. Patent 9,918,948 in March 2018
- · 2 patent applications pending



TNX-102 SL: Sublingual Formulation is Designed for Bedtime Administration



10

- · Innovation by design with patent protected CBP/mannitol eutectic
- Rapid systemic exposure
- · Increases bioavailability during sleep
- Avoids first-pass metabolism
- · Lowers exposure to long-lived active major metabolite, norcyclobenzaprine (norCBP)

CBP undergoes extensive first-pass hepatic metabolism when orally ingested

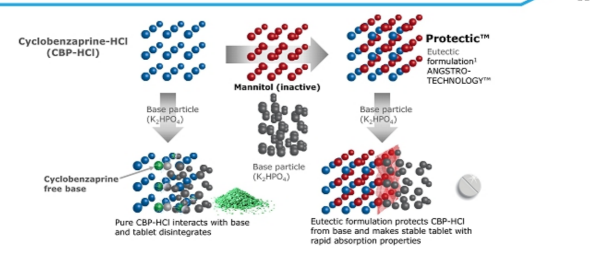
- Active major metabolite, norCBP¹
 - Long half-life (~72 hours)
 - Less selective for target receptors (5-HT_{2A}, α₁-adrenergic, histamine H₁)
 - More selective for norepinephrine transporter and muscarinic M₁

TNX-102 SL 505(b)(2) NDA approval can rely on the safety of the reference listed drug (AMRIX[®])²

¹ Daugherty et al., Abstract 728, Society of Biological Psychiatry 70th Annual Scientific Carvention, May 14-16, 2015, Toronto Ontario, Canada ² FDA Minutes (November 26, 2018) © 2019 Tonix Pharmaceuticals Holding Corp.



Proprietary Cyclobenzaprine Hydrochloride Eutectic Mixture Stabilizes Sublingual Tablet Formulation



¹U.S. Patent issued May 2, 2017



Tonmya: Hypothesized Novel Mechanism Targets Sleep Quality for Recovery from PTSD



PTSD is a disorder of recovery

- · Most people exposed to extreme trauma recover over a few weeks
- In PTSD, recovery process impeded due to insufficient sleep-dependent memory processing^{1,2}

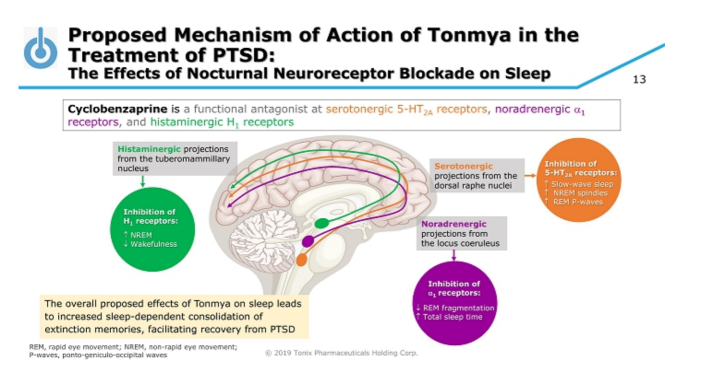
Memory processing is essential to recovery

 Vulnerability to memory intrusions and trauma triggers remains if no consolidation of new learning (extinction)

Tonmya targets sleep quality³

 The active ingredient in Tonmya, cyclobenzaprine, interacts with receptors that regulate sleep quality: strongly binds and potently blocks 5-HT_{2A}, α₁-adrenergic and histamine H₁ receptors, permissive to sleep-dependent recovery processes

¹Straus LD, Acheson DT, Risbrough VB, Drummond SPA. Sleep Deprivation Disrupts Recall of Conditioned Fear Extinction. Biol Psychiatry Cogn Neurosci Neuroimaging. 2017; 2(2):123-129. "Murkar ALA, De Koninck J. Consolidative mechanisms of emotional processing in REM sleep and PTSD. Sleep Med Rev. 2018; 41:173-184. "Daugherty et al., Abstract 728, Society of Biological Psychiatry 70th Annual Scientific Convention, May 14-16, 2015, Toronto Ontario, Canada



6

Proposed Mechanism of Action of Tonmya in the Treatment of PTSD: Focus on Nocturnal 5-HT_{2A} Receptor Blockade in REM



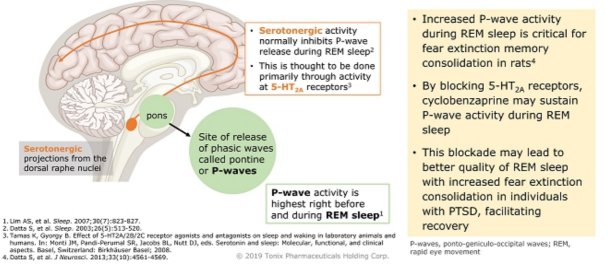
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- Extinction learning is critical to recovery from trauma, and such new learning is consolidated (moving from labile short term to established long term memory) during particular stages of sleep^{1,2}
- Recent rodent research shows how particular brain wave patterns during REM sleep, known as "P-waves" are critical to extinction consolidation³
- 5-HT activation of pontine brainstem region richly expressing 5-HT $_{\rm 2A}$ receptors inhibits P-wave generation during REM 4
- Nocturnal blockage of 5-HT_{2A} receptors may restore extinction consolidation by inhibition of errant 5-HT stimulation during REM (see model in next 2 slides)

1. Pace-Schott, et al. Biology of Mood & Anxiety Disorders. 2015;5(3):1-19. 2. Straus et al. Biol Psych: CNNI. 2017;2(2):123-129. 3. Datts 5, et al. J Neurosc. 2013;33(10):4561-4569. 4. Datts 5, et al. Sitep. 2003;26(5):513-520. © 2019 Tonix Pharmaceuticals Holding Corp.



Fear Extinction Memory Consolidation: The Proposed Role of P-Waves, REM Sleep, and Serotonergic Neuroreceptor Activity



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P-waves, ponto-geniculo-occipital waves; REM, rapid eye movement

15



Overview of Posttraumatic Stress Disorder (PTSD)

16

PSTD is a chronic disabling disorder in response to experiencing traumatic event(s)

Symptoms of PTSD fall into four clusters:

- 1. Intrusion (aversive memories, nightmares, flashbacks)
- 2. Avoidance (avoiding persons, places or situations)
- 3. Mood/cognitions (memory block, emotional numbing, detachment from others)
- 4. Hyperarousal (anxiety, agitation & sleep disturbance)

Diagnosis, symptom severity, as well as treatment effect, is determined by CAPS-5*

- · Recognized as the standard for rating PTSD severity in clinical trials
- · Takes into account all four symptom clusters
- · Higher Total CAPS-5 score reflects more severe PTSD symptoms

* Clinician-administered PTSD scale for Diagnostic Statistical Manual version 5 (DSM-5)



Consequences:

Impaired daily function and substantial interference with work and social interactions

17

- · Reckless or destructive behavior
- · Increased health care utilization and greater medical morbidity

PTSD as a risk factor for:

- Depression
- · Alcohol or substance abuse
- Absenteeism/unemployment
- Homelessness
- Violent acts
- · Suicidal thoughts and suicide



PTSD: U.S. Prevalence and Index Traumas



PTSD is a chronic response to traumatic event(s)

· A majority of people will experience a traumatic event at some point in their lifetime1

20% of women and 8% of men in the U.S. who experience significant trauma develop PTSD¹

Adult Civilians:

- 6.1% (14.4 million adults in the U.S.)2 Lifetime prevalence:
 - Persistent >1/3 fail to recover, even after several years following the trauma²

<u>Twelve month prevalence</u>: U.S. 4.7% (12 million adults)²

EU 2.3% (~10.0 million adults)³

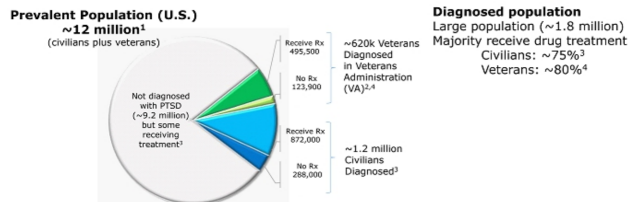
Most common forms of trauma¹

- · Witnessing someone being badly injured or killed
- Natural disaster
- · Life-threatening accident
- Sexual or physical assault

¹ Kessler et al., Arch Gen Psychiatry 1995; 52:1048
 ² Goldstein et al., 2016 (adjusted for 2019)
 ³ The European Union Market Potential for a New PTSD Drug. Prepared for Tonix Pharmaceuticals by Procela Consultants Ltd, September 2016

PTSD Prevalence and Market Characteristics





¹ Goldstein et al., 2016 (adjusted for 2019)
 ² Veterans: VA/DOD Clinical Practice Guidelines for the Managements of PTSD and Acute Stress Disorder, 2017, page 15 (619,493 vets diagnosed with PTSD in VA for 2016)
 ³ IMS Consulting, Market Sizing & Treatment Dynamics: Post-Traumatic Stress Disorder (PTSD) Patients", 2016
 ⁴ Bernardy et al., 2012 (80% of veterans diagnosed with PTSD had at least one medication from the Clinical Practice Guidelines)

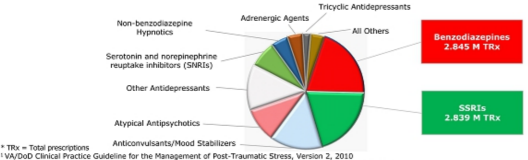


Market highly fragmented, with benzodiazepines widely prescribed (but not indicated)¹

20

- Multiple medications per patient (or "Polypharmacy") is the norm
- Approximately 55% of patients receive a benzodiazepine, and 53% receive a selective serotonin reuptake inhibitor (SSRI)
- · SSRIs are the only FDA-approved drug class

Estimated PTSD Market Volume (Civilian Population Only) ~14.1 million TRx*2



¹ VA/DoD Clinical Practice Guideline for the Management of Post-Traumatic Stress, Version 2, 2010
 ² IMS Consulting, Market Sizing & Treatment Dynamics: "Post-Traumatic Stress Disorder (PTSD) Patients", 2016
 (8) 2019 Tonix Pharmaceuticals Helding Corp.



FDA-approved SSRIs, paroxetine and sertraline, are indicated as a treatment for PTSD

21

- · Neither drug has shown efficacy in military-related PTSD
- · Majority of male PTSD patients unresponsive or intolerant to current treatments
- Side effects relating to sexual dysfunction, sleep disturbance and weight gain are commonly reported

Characteristics of an ideal drug therapy that would be compatible and complementary with behavioral therapy

- Lack of retrograde amnesia (e.g., unlike off-label use of benzodiazepines and nonbenzodiazepines)
- · Lack of interference on sleep (e.g., unlike approved SSRIs)

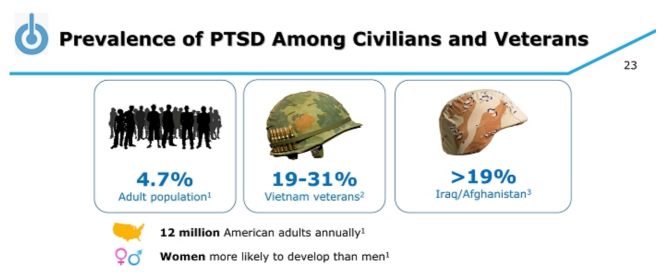
Tonmya is being investigated in both military and civilian PTSD and is expected to be indicated as a "treatment for PTSD"



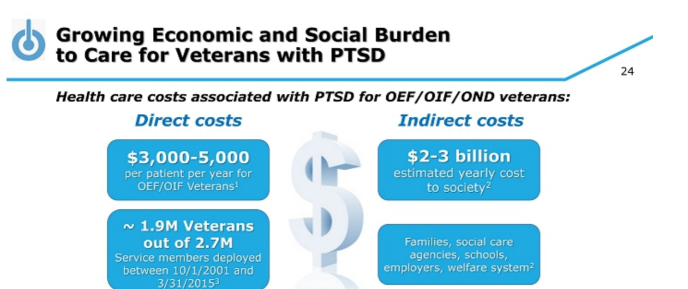


Military-related PTSD not well-served by existing FDA-approved therapies

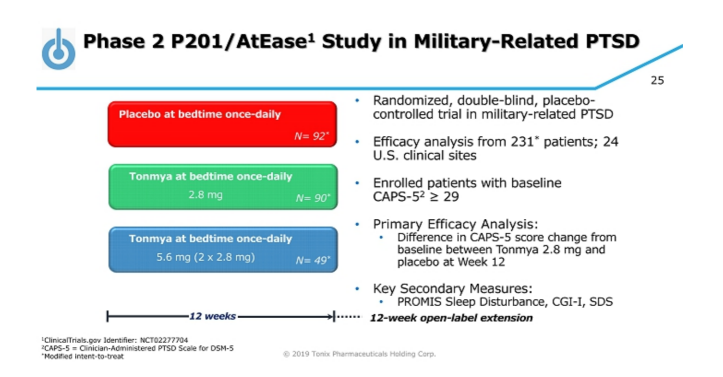
	No clear treatment response observed in U.S. military population
	Sertraline: failed to show efficacy in a large multicenter trial in U.S. military (placebo numerically better) ¹ Paroxetine: no large trials conducted with predominantly military trauma
•	Inconsistent treatment response observed in males
	Sertraline: FDA-conducted post-hoc analysis concluded no effect for male civilian subgroup ² Paroxetine: no sex-related difference in treatment outcomes ³
	Important tolerability issues with SSRIs in this population
	Sexual dysfunction ^{2,3} Insomnia ^{2,3} SSRI withdrawal syndrome ⁴
² Zoloft Package Ins ³ Paxil Package Inse	



³Goldstein et al., 2016 (adjusted for 2019); ²Norris, PTSD Res Quar. 2013; ³Analysis of VA Health Care Utilization among Operation Enduring Freedom, Operation Iragi Freedom, and Operation New Dawn Veterans, from 1st Qtr FY 2002 through 2nd Qtr FY 2015, Washington, DC; Among 1.9M separated OEF/OIF/OND veterans, 1.2M have obtained VA healthcare; 685k evaluated by VA with possible mental disorder, and 379k diagnosed with PTSD.



¹ CBO Report 2012; ² Tanielan, Invisible Wounds of War. 2005; ¹ Analysis of VA Health Care Utilization among Operation Enduring Freedom, Operation Iraqi Freedom, and Operation New Dawn Veterans, from 1st Qtr FY 2002 through 2nd Qtr FY 2015, Washington, DC; OEF/OIF/OND, Operations Enduring Freedom, Iraqi Freedom and New Dawn. © 2019 Tonix Pharmaceuticals Holding Corp.







P201 was a large adequate well-controlled Phase 2 study in militaryrelated PTSD

- Primary endpoint (Week 12 CAPS-5) did not separate from placebo for TNX-102 SL 2.8 mg
- · No safety or tolerability issue discovered
- Retrospective analyses showed TNX-102 SL 5.6 mg had a strong signal of treatment effect at Week 12 CAPS-5 (P=0.053) and CGI-I (P=0.041) scores
- Retrospective analyses suggested CAPS-5 ≥ 33 enrollment criteria for Phase 3



P201/AtEase Study – Summary of Primary and Secondary Analyses (Week 12)



Assessment	Domain	Analysis	p-Values	
			2.8 mg (N=90)	5.6 mg (N=49)
CAPS-5	Total	MMRM (Primary Analysis)	0.259^	0.053
	Total	MMRM with Multiple Imputation	0.211	0.031*
	Total	MMRM w/ Hybrid LOCF/BOCF	0.172	0.037*
	Total	MMRM with Multiple Imputatio MMRM w/ Hybrid LOCF/BOCF ANCOVA I & Reactivity cluster (E) MMRM tem (E6) MMRM erated Startle item (E4) MMRM iders Logistic Regression icore MMRM icolo item MMRM	0.090	0.038*
CAPS-5 clusters/items	Arousal & Reactivity cluster (E)	MMRM	0.141	0.048*
	Sleep item (E6)	MMRM	0.185	0.010*
	Exaggerated Startle item (E4)	MMRM	0.336	0.015*
CGI-I	Responders	Logistic Regression	0.240	0.041*
PGIC	Mean score	MMRM	0.075	0.035*
Sheehan Disability Scale	Work/school item	MMRM	0.123	0.050*
	Social/leisure item	MMRM	0.198	0.031*
MMRM, mixed model repeated	ried forward; CGI-I, Clinical Global : measures; PGIC, Patient Global Imp significant comparing Tonmya 2.8 m		CF, last observation ca	arried forward;

P301/HONOR¹ Study – Evidence of Efficacy at Week 4 Discontinued Due to High Placebo Response at Week 12

General study characteristics:

Randomized, double-blind, placebo-controlled, adaptive design, planned 550 military-related PTSD participants with baseline CAPS- $5^2 \ge 33$ in approximately 40 U.S. sites

Primary endpoint CAPS-5²:

 Mean change from baseline at Week 12 (Tonmya 5.6 mg vs. placebo)

28

Tonmya once-daily at bedtime5.6 mg (2 x 2.8 mg tablets) $N= 125^*$ Placebo once-daily at bedtimeN= 127*

– 12 weeks -

Unblinded interim analysis at 274 randomized participants (mITT* N= 252)

- Study stopped due to not meeting a pre-specified study continuation threshold at Week 12
- Participants discontinued in HONOR or 12-week open-label extension (OLE) studies can enroll in the 40-week OLE study

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¹ClinicalTrials.gov Identifier: NCT03062540 ²CAPS-5 = Clinician-Administered PTSD Scale for DSM-5 *Modified intent-to-treat



P301/HONOR Study- Primary Analysis in mITT Population

	Placebo N=127		TNX-102 S		
Visit			N=1		
Statistic	CAPS-5 Value	MCFB	CAPS-5 Value	MCFB	Difference
Week 4					
LS Mean (SE)	31.0 (1.62)	-11.2 (1.62)	27.5 (1.73)	-14.7 (1.73)	-3.6 (1.51)
95% CI	(27.8,34.2)	(-14.4,-8.0)	(24.1,30.9)	$\{-18.1, -11.4\}$	(-6.5,-0.6)
p-value					0.019
Week 8					
LS Mean (SE)	29.4 (1.76)	-12.8 (1.76)	27.6 (1.86)	-14.6 (1.86)	-1.8 (1.77)
95% CI	(25.9,32.8)	(-16.3,-9.4)	(24.0,31.3)	$\{-18.2, -10.9\}$	(-5.2,1.7)
p-value					0.321
Week 12					
LS Mean (SE)	28.0 (1.80)	-14.2 (1.80)	27.0 (1.90)	-15.2 (1.90)	-1.0 (1.88)
95% CI	(24.5,31.5)	(-17.7,-10.7)	(23.3,30.8)	$\{-18.9, -11.4\}$	(-4.7,2.7)
p-value					0.602

29

MMRM with Multiple Imputation

In P301 study both TNX-102 SL and placebo-treated groups improved but the greater improvement on TNX-102 SL compared with placebo diminished over time

TNX-102 SL did not separate from placebo at primary endpoint

LS Mean (SE) = Least Squares Mean (Standard Error) CI = Confidence Interval MCFB = Mean Change From Baseline



TNX-102 SL: Sublingual Formulation is Designed for Bedtime Administration



with transmucosal absorption

- · Innovation by design with patent protected CBP/mannitol eutectic
- Rapid systemic exposure
- · Increases bioavailability during sleep
- Avoids first-pass metabolism
- Lowers exposure to long-lived active major metabolite, norcyclobenzaprine (norCBP)

CBP undergoes extensive first-pass hepatic metabolism when orally ingested

- Active major metabolite, norCBP¹
 - Long half-life (~72 hours)
 - Less selective for target receptors (5-HT_{2A}, α₁-adrenergic, histamine H₁)
 - · More selective for norepinephrine transporter

TNX-102 SL 505(b)(2) NDA approval can rely on the safety of the reference listed drug (AMRIX[®])²

¹ Daugherty et al., Abstract 728, Society of Biological Psychiatry 70th Annual Scientific Convention, May 14-16, 2015, Toronto Ontario, Canada
² FDA Meeting Minutes (November 26, 2018)
^(a) 2019 Tonix Pharmaceuticals Holding Corp.



Differences Between P201/AtEase and P301/HONOR Studies Design

Categories	P201	P301
No. of US Sites Randomizing ≥ 1	24	43
No. of Treatment Arms	3	2
Baseline Entry CAPS-5 Threshold	≥ 29	≥33
Range of Includable Ages, years	18-65	18-75
Depression Rating Scale Employed	MADRS	BDI-II
Minimum Time Since No TFT	1 month	3 months
Primary Endpoint Analytic Method	MMRM	MMRM with MI
No. of In-Clinic Study Visits	9	5
No. of CAPS-5 Administrations	6	5
Key Secondary Endpoints	CGI-I, SDS, PROMIS SD	CGI-I, SDS

31

Phase 2 and 3 studies were very similar – both studied military related PTSD at multiple sites in the US

CAPS-5 ≥ 33 entry criteria used in Phase 3

BDI-III= Beck Depression Inventory-II; CGI-I=Clinical Global Impression – Improvement; MI= multiple imputation; MMRM=mixed model repeated measures; MADRS=Montgomery-Åsberg Depression Rating Scale; PROMIS SD=Patient-Reported Outcomes Measurement Information System – Sleep Disturbance; SDS=Sheehan Disability Scale; TFT=trauma-focused therapy © 2019 Tonix Pharmaceuticals Holding Corp.



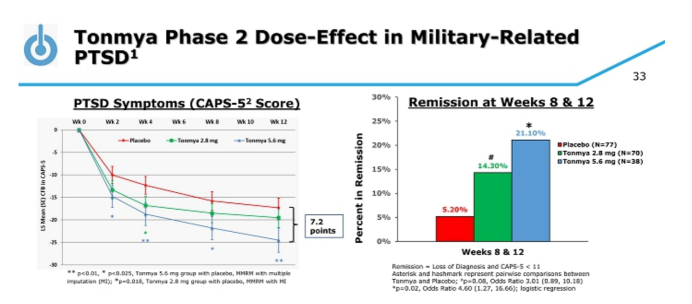
P201/AtEase and P301/HONOR Demographics and Characteristics

32

	P201			P301	
Variable	Placebo N=92	TNX 2.8 mg N=90	TNX 5.6 mg N=49	Placebo N=127	TNX 5.6 mg N=125
Females, %	6.50%	6.70%	8.20%	13.40%	8.00%
Age, yrs. (SD)	32.0	34.5	34.8	35.5	35.9
Body Mass Index, kg/m ²	28.9	29.0	29.0	29.3	29.9
Employment (current), %	58.7%	62.2%	67.3%	63.0%	55.2%
Unable to work due to PTSD, %	9.8%	11.1%	14.3%	12.6%	16.8%
Active Duty/Reservists/Veterans, No.	8/4/79	9/5/71	5/7/37	17/0/110	9/0/116
Time since trauma, mean years	7.1	7.3	6.2	9.2	9.2
Time since trauma, median years	7.0	7.2	6.0	9.3	9.5
Combat index trauma, %	80.4%	85.6%	93.8%	77.2%	83.2%
Number of deployments	2.2	2.3	2.6	3.0	2.6
Baseline CAPS-5 Scores	39.5	39.5	39.3	42.4	42.0
Baseline BDI-II Scores	NA	NA	NA	23.0	25.6
Baseline MADRS Scores	17.3	17.6	16.1	NA	NA

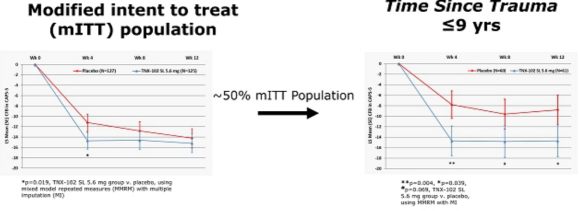
The striking difference between P201 and P301 was time since trauma

 Phase 2 P201 study recruited many participants from the surge in Iraq who were mostly <9 years since trauma



³Completed Phase 2 P201/AtEase study: Retrospective analysis of Tonmya 5.6 mg on CAPS-5 ≥33 (high-moderate) subgroup. Primary analysis of P201/AtEase was on Tonmya 2.8 mg in participants with entry CAPS-5 ≥29, moderate PTSD severity. ³Clinician administered PTSD Scale for DSM-5

Primary Outcome (CAPS-5) in Phase 3 (mITT) and ≤9 Years Time Since Trauma (TST) Subgroups Phase 3 P301/HONOR Study¹ Modified intent to treat Time Since Trauma



¹Phase 3 P301/HONOR study: stopped in July 2018. Separation on primary endpoint did not cross pre-specified study continuation threshold at Week 12 in the interim analysis at ~50% randomization; no safety or tolerability issues discovered. (2) 2019 Tonix Pharmaceuticals Holding Corp.



Retrospective Comparison of Time Since Trauma in P201/AtEase versus P301/HONOR (Tonmya 5.6 mg Groups)

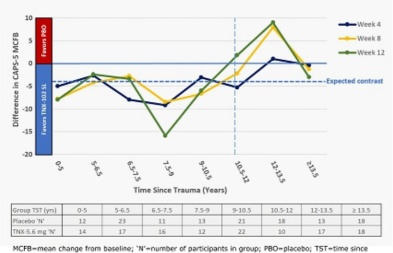
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P301 study was initiated approximately two years later than Phase 2 P201

 The median time since trauma in Phase 3 was 9.5 years compared to the median time since trauma in Phase 2 of 6.0 years for TNX-102 SL 5.6 mg treated groups

CAPS-5 Mean Change from Baseline Difference from Placebo of Tonmya 5.6 mg in TST Subgroups in P301¹



trauma (© 2019 Tonix Pharmaceuticals Holding Corp

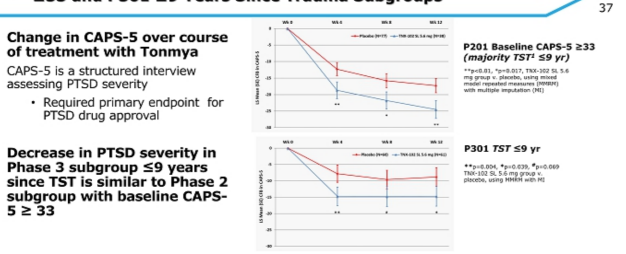
 The mITT population was divided into subgroups based on TST (1.5-2 years each as well as 0-5 years and ≥13.5 years subgroups)

36

- Graph shows the CAPS-5 differences in MCFB between TNX 5.6 mg and PBO for Weeks 4, 8, and 12 post-baseline timepoints
- and 12 post-baseline timepoints
 "Expected contrast" horizontal dashed line indicates observed effect from Phase 2 P201 study
- For TST <10.5 years groups, TNX 5.6 mg showed good separation from PBO (left side of vertical dashed 10.5 year line)
- For TST >10.5 years groups, separation of TNX 5.6 mg from PBO was either small or worked in the favor of PBO (right side of vertical dashed 10.5 year line)

¹Time Since Trauma in PTSD: Phase 3 Multi-Center, Double-Bilind, Placebo-Controlled Trial of TNX-102 SL, a Sublingual Formulation of Cyclobenzaprine, in Military-Related PTSD (Study TNX-CY-P301) Presented at CNS Summit in Boca Raton, FL November 1-4, 2018 and abstract published in Innovations in Clinical Neuroscience, November-December 2018;15(11-12, suppl):S10. https://content.equisolve.net/tonixpharma/media/1d0c405 5b2863fc74e1ef45/9ddaf42b.pdf

PTSD Treatment Response to Tonmya in Phase 2 and Phase 3 Studies: Retrospective Analyses of P201 Entry CAPS-5 ≥33 and P301 ≤9 Years Since Trauma Subgroups



¹Time since trauma; ²Majority of P201 participants were ≤9 years since trauma and ~80% of P201 participants and all of P301 participants were ≥33 CAPS-5 at baseline

6

Sustained Remission in Phase 2 and Phase 3 Studies: Retrospective Analyses of P201 Entry CAPS-5 ≥33 and P301 ≤9 Years Since Trauma Subgroups

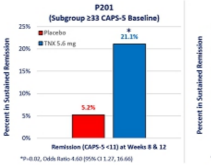
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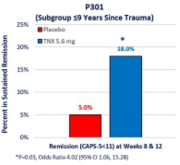
Remission is a clinical state that is essentially asymptomatic

In order to confirm remission:

 Determined rates of participants who met remission status at both Week 8 and Week 12

Rate of remission in ≤9 years since trauma group in P301 is similar to baseline CAPS-5 ≥ 33 group in P201¹





¹Majority of P201 participants were ≤ 9 years since trauma and ~80% of P201 participants and all of P301 participants were ≥ 33 CAPS-5 at baseline



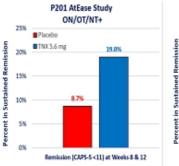
Sustained Remission in P201/AtEase Study Retrospective Analyses of Phase 2 Subgroups with and without Oral AE's (ON/OT/NT)

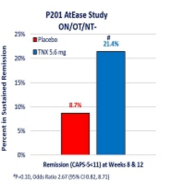
Oral numbness (ON), oral tingling (OT) and noticeable taste (NT) are local administration site reactions that are potentially unblinding

 Subgroups with and without ON/OT/NT were studied in participants who met remission status at *both* Week 8 and Week 12

Similar rates of remission were observed in participants in P201 with and without oral AE's

 Unblinding was unlikely to account for treatment effect





39



Betrospective Analyses of ≤9 Years Since Trauma Subgroup on Primary and Secondary Endpoints in P301/HONOR Study

			P301 mITT			P301 ≤9 Year Subgroup					
			PBO (N=127) v. TNX-5.6 (N=125)				PBO (N=60) v. TNX-5.6 (N=61)				
			Wee	ek 4	Week 12		Week 4		Week 12		
Measure Analysis		LSMD	p-value	LSMD	p-value	LSMD	p-value	LSMD	p-value		
1°	CAPS-5	MMRM/MI	-3.6	0.019	-1.0	0.602	-6.9	0.004	-5.9	0.039	
2°s	CGI-I	MMRM	-0.3	0.015	-0.1	0.403	-0.6	0.002	-0.5	0.021	
	SDS	MMRM	-0.2	0.785	-1.6	0.101	-1.8	0.167	-4.3	0.007	
	PGIC	MMRM	-0.2	0.238	-0.3	0.020	-0.4	0.045	-0.6	0.007	
	PROMIS SD	MMRM	-3.1	0.015	-2.7	0.082	-4.5	0.029	-5.0	0.042	
	BDI-II	MMRM	-1.1	0.330	-1.4	0.255	-5.2	0.008	-6.6	0.001	

BOLDED p-values are all p<0.05; BDI-II=Beck Depression CAPS-5=Clinician-Administered PTSD Scale for DSM-5; CGI-I=Clinical Global Impression - Improvement scale; mITT=modified Intent-to-Treat sample; MNRM=mixed model repeated measures analysis; MI=multiple imputation; PGIC=Patient Global Impression of Change scale; RROMIS SD=Patient-Reported Outcome Measurement Information System Sleep Disturbance instrument (short form 8a); P80=placebo; SDS=Sheehan Disability Scale; TNX-5.6=TNX-102 SL 5.6 mg; yrs=years; 1°=primary; 2°s=secondaries

Secondary endpoints also showed strong treatment effects in ≤9 yrs TST

· Support CAPS-5 results and similar to Phase 2 P201 Study results



Retrospective Analyses of ≤9 Years Since Trauma Subgroup on Key Secondary Endpoints in P301/HONOR Study

41

			P301	miTT		P301 ≤9 Year Subgroup					
		PBO (N=	PBO (N=127) v. TNX 5.6 mg (N=125)				PBO (N=60) v. TNX 5.6 mg (N=61)				
		We	ek 4	Week 12		Week 4		Week 12			
	Analysis	LSMD	p-value	LSMD	p-value	LSMD	p-value	LSMD	p-value		
CGI-I	MMRM	-0.3	0.015	-0.1	0.403	-0.6	0.002	-0.5	0.021		
PGIC	MMRM	-0.2	0.238	-0.3	0.020	-0.4	0.045	-0.6	0.007		
SDS	MMRM	-0.2	0.785	-1.6	0.101	-1.8	0.167	-4.3	0.007		
PROMIS SD	MMRM	-3.1	0.015	-2.7	0.082	-4.5	0.029	-5.0	0.042		

Key secondary endpoints showed strong treatment effects

- · CGI-I, PGIC and PROMIS SD were pre-specified secondary analyses
- · Supports CAPS-5 results and similar to Phase 2 P201 Study results

CGI-I=Clinical Global Impressions - Improvement scale PGIC, Patient Global Impression of Change scale PROMIS SD=Patient-Reported Outcome Neasures Information System Sleep Disturbance SDS=Sheehan Disability Scale LSMD = Lesst Squares Mean Difference



Adverse Events (AEs) in P201/AtEase and P301/HONOR Studies

	P201			P301	
Category of Adverse Reaction Preferred Term	Placebo (N=94)	TNX 2.8 mg (N=93)	TNX 5.6 mg (N=50)	Placebo (N=134)	TNX 5.6 mg (N=134)
Systemic Adverse Events**					
Somnolence	6.4%	11.8%	16.0%	9.0%	15.7%
Dry mouth	10.6%	4.3%	16.0%		
Headache	4.3%	5.4%	12.0%		
Insomnia	8.5%	7.5%	6.0%		
Sedation	1.1%	2.2%	12.0%		
Local Administration Site Reactions*					
Hypoaesthesia oral	2.1%	38.7%	36.0%	1.5%	37.3%
Paraesthesia oral	3.2%	16.1%	4.0%	0.7%	9.7%
Glossodynia	1.1%	3.2%	6.0%		
Product Taste Abnormal				3.0%	11.9%

42

²only adverse events (AEs) are listed that are at a rate of \geq 5% in any TNX-treated group *no values in a row for either study means the AE in the active group(s) in that study was at a rate of <5%

No serious or unexpected AEs in P201 or P301 related to Tonmya

- Systemic AEs comparable between studies and also consistent with those described in approved oral cyclobenzaprine product labeling
- Severity and incidence of oral hypoesthesia (oral numbness) are not dose related and similar in both studies
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Time Since Trauma – Review of Published Studies

43

/w

Published studies of prazosin Time Since Trauma suggested effects in military-PTSD 0 1 2 5 4 5 6 7 8 9 10 11 12 15 14 15 16 17 18 19 20 Time Singe Trauma (Neuro) prior to 9 years PROT-OF Years P301 >9 Years Loss of treatment effect >9 years P20 Paroxetine and sertraline studies supporting FDA approval were Plaquetine* conducted on PTSD > 9 years · SSRIs have a benefit long after pertrauma ¹Martenyi et al. J Clin Psychiatry 2002;63:199-205. ²Friedman et al. J Clin Psychiatry 2007;68:711-720. ³Raskind et al. ACM 2010;378:507-517. ⁴Raskind et al. ACM J Psychiatry 2012;709:1003-1010. ⁵Shalev et al. ACM Gen Psychiatry 2012;69:166-176. ⁵Davidson et al. ACM Gen Psychiatry 2001;58:485-492. ³Brady et al. JAMA 2000;283:1837-1844. ⁴Marshall et al. ACM J Psychiatry 2001;58:1882-1988. ⁵Tucker et al. J Clin Psychiatry 2001;62:860-868. 0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 EscR=escRalopram



Time Since Trauma – Remitting and Persistent Phases of PTSD

Kessler et al¹ studied remission in PTSD with and without therapy

- · Identified remitting and persistent phase of PTSD with transition at approximately 6 years post trauma
- Supported by other studies2-6 ٠

 ¹Kessler et al. Arch Gen Psychiatry 1995;52:1048-1060.

 ²Armenta et al. BMC Psychiatry 2018;18:48.

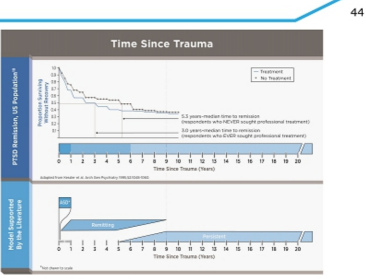
 ³Salatzer-Levy et al. PLOS OWE 2013;8:e70084.

 ³Perkoning et al. Arch Schröder 2005;162:1320-1327.

 ³Santiago et al. PLOS OWE 2013;8:e59236.

 ³Davidson & Connor. Eur Neuropsychopharmacol 2001;11(Supp3):5148-5149

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Response to Tonmya for Female Participants in P301/HONOR Study¹



Females made up only 11% of the P301/HONOR study mITT population

Difference in mean change from baseline in CAPS-5 in females between placebo (N=17) and Tonmya 5.6 mg (N=10) was:

- At 4 weeks -11.5 points
- · At 12 weeks -9.1 points

Indicates substantial separation from placebo in the small number of female participants

Predicts therapeutic response to Tonmya 5.6 mg likely in mixed civilian and military PTSD population to be studied in upcoming P302/RECOVERY trial

Civilian PTSD population tends to be about 2/3 female

¹Time Since Trauma in PTSD: Phase 3 Multi-Center, Double-Blind, Placebo-Controlled Trial of TNX-102 SL, a Sublingual Formulation of Cyclobenzaprine, in Military-Related PTSD (Study TNX-CY-P301) Presented at CNS Summit in Boca Raton, FL November 1-4, 2018 and abstract published in Innovations in Clinical Neuroscience, November-December 2018;15(11-12,suppl):S10.<u>https://content.eguisolve.net/tonixpharma/media/1d0c4055b2863fc74e1ef45f9ddaf42b.pdf</u>

Response to Tonmya for Non-Combat Traumas in P301/HONOR Study in ≤9 Years Time Since Trauma Subgroup¹

Non-combat traumas studied are similar to traumas experienced in civilian populations with PTSD

To determine the therapeutic effects of Tonmya 5.6 mg in a mixed civilian and military population, difference in MCFB in CAPS-5 was assessed in non-combat traumas in \leq 9 years TST subgroup (placebo N=14, Tonmya 5.6 mg N=10):

46

- At 4 weeks -4.8 points
- At 12 weeks -4.4 points

Non-combat traumas treated with Tonmya 5.6 mg showed clinically meaningful separation from placebo at Weeks 4 and 12, suggesting a mixed civilian and military sample within 9 years of index trauma may show a therapeutic response to Tonmya

¹Time Since Trauma in PTSD: Phase 3 Multi-Center, Double-Blind, Placebo-Controlled Trial of TNX-102 SL, a Sublingual Formulation of Cyclobenzaprine, in Military-Related PTSD (Study TNX-CY-P301) Presented at CNS Summit in Boca Raton, FL November, 2018; and abstract published in Innovations in Clinical Neuroscience, November-December 2018;15(11-12,suppl):S10.https://content.equisolve.net/tonixpharma/media/1d0c4055b2863fc74e1ef45f9ddaf42b.pdf

CAPS-5=Clinician-Administered PTSD Scale for DSM-5; MCFB=mean change from baseline; mITT=modified Intent-to-Treat sample; TST=time since trauma
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Summary of Clinical Experience with Tonmya/ TNX-102 SL in PTSD

Median time since trauma (TST) in TNX-102 SL 5.6 mg group in the P301/HONOR study (9.5 years) was longer than P201/AtEase study (6 years)

47

- · Both studied military-related PTSD
- · Time has passed since the surge in Iraq

In retrospective analysis, the \leq 9 year subgroup of P301 study had similar results as the P201 study (primary and secondary)

- · TST is important in placebo-controlled clinical study
- Potential enrichment in ≤ 9 years TST subgroup for treatment responders

The ≤ 9 year subgroup of P301 may be enriched for "Remitting Phase" of PTSD¹⁻⁴

· Expect remitting phase of PTSD is more amenable to drug studies

Results from retrospective analyses lead to improved Phase 3 study design

Kessler et al. Arch Gen Psychiatry 1995;52:1048-1060. Varmenta et al. BMC Psychiatry 2018;18:48. TGalatzer-Lavy et al. PLOS ONE 2013;8:e70084. Perkonigg et al. Am J Psychiatry 2005;162:1320-1327.



New Phase 3 P302/RECOVERY Study – Expected to Start 1Q 2019



General study characteristics: Primary endpoint: Randomized, double-blind, placebo-controlled study with CAPS-5¹ mean change from baseline at Week 4 (Tonmya 5.6 mg vs. placebo) . baseline CAPS-5¹ ≥ 33 in approximately 30 U.S. sites · Enrollment restricted to study participants with PTSD who experienced an index trauma \leq 9 years from the date of Key Secondary endpoints include: screening · CAPS-5 mean change from baseline at Week 12 (Tonmya 5.6 mg · Both civilian and military-related PTSD to be included vs. placebo) · Change from baseline Clinical Global Impression - Severity scale Tonmya once-daily at bedtime · Change from baseline Sheehan Disability Scale total score Potential pivotal efficacy study to support NDA approval Placebo once-daily at bedtime 12 weeks -≁ Primary endpoint CAPS-51 at Week 4 CAPS-5 = Clinician-Administered PTSD Scale for DSM-5 @ 2019 Tonix Pharmaceuticals Holding Corp.



Late-Stage PTSD Drug Candidates

Tonmya

 Phase 3 development focused on military-related and civilian PTSD; showed activity in treatment of military-related PTSD in large multi-center trials 49

MDMA-assisted psychotherapy

 Breakthrough therapy that is Phase 3-ready; showed activity in a Phase 2 study of PTSD; enrolling in Phase 3

Other drugs currently (or recently) in Phase 2 development

- Rexulti® (brexpiprazole) Otsuka/Lundbeck; atypical antipsychotic; positive clinical results from Phase 2 study reported in November 2018 for brexpiprazole, when used in combination with an approved PTSD medication, sertraline, but not as monotherapy
- NYX-783 Aptinyx; NMDA receptor modulator (enrolling for 8-week Phase 2 study of 144 patients using 50 mg either once daily or once weekly)
- BNC-201 Bionomics; nicotinic receptor modulator (program planned to resume after reformulation)





Tonix is exploring a variety of options to commercialize TNX-102 SL, including commercializing on our own or partnering all or some indications in specific regions of the world

Tonix has participated in numerous partnering meetings

Commercial Considerations:

- Primary physician audience is well defined: psychiatrists (~30,000 in U.S.)
 - · Small specialty sales force sufficient for coverage
- Primary market research with psychiatrists indicate strong interest in new therapeutic options



TNX-102 SL – Multi-Functional Mechanism Involves Antagonism at 3 Neuronal Receptors

51

Active ingredient, cyclobenzaprine, interacts with 3 receptors

- Antagonist at 5-HT_{2A} receptors
 - Similar activity to trazodone and Nuplazid[®] (pimivanserin)
- Antagonist at α₁-adrenergic receptor
 - Similar activity to prazosin
- Antagonist at histamine H₁ receptors
 - · Similar activity to Benadryl® (diphenhydramine) and hydroxyzine

Multi-functional activity suggests potential for other indications

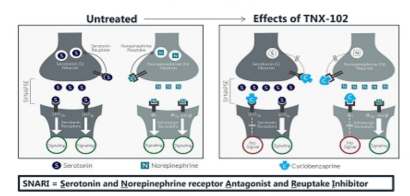
- TNX-102 SL was developed for the management of fibromyalgia (Phase 3)
- Sleep quality is a problem in other conditions





Cyclobenzaprine is a multi-functional drug - SNARI

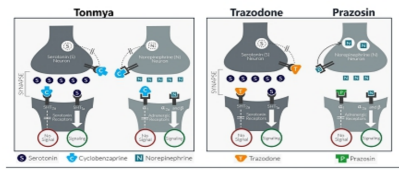
- inhibits serotonin and norepinephrine reuptake
- inhibits serotonin and norepinephrine reuptane blocks serotonin 5-HT_2A and norepinephrine α_1 receptors







- Trazodone (disordered sleep), prazosin (night terrors) Trazodone inhibits serotonin 5HT_{2A} receptors and serotonin reuptake (SARI)
 - Prazosin blocks norepinephrine α₁ receptors



SARI - Serotonin Receptor Antagonist & Beuptake Inhibitor (Stahl SM, CNS Spectrums, 2009;14:536).



Role of sleep disturbance more established in common psychiatric and neurological/pain disorders

- · Recognized as a core symptom of many of these disorders
- Traditional sleep medications, which increase sleep quantity, may not provide benefit (benzodiazepines in major depression) or are contraindicated (benzodiazepines in PTSD)

Psychiatric Disorders

- Stress Disorders (PTSD)
- Mood Disorders
- Anxiety Disorders

Psychiatric Symptoms of Neurological Disorders

- Agitation in Alzheimer's
 Psychosis in Parkinson's, Alzheimer's and other
- dementias

Chronic Pain States

 Chronic wide-spread pain (fibromyalgia)

54

Osteoarthritis

Growing recognition that there are many disorders where sleep disturbances may have a role in the pathophysiology (cardiovascular, metabolic, neurologic)

· Homeostatic role of sleep quality in several disorders



TNX-102 SL – Bedtime Treatment for Multiple Potential Indications



Management of Fibromyalgia (FM) – chronic pain condition

- TNX-102 SL studied at low dose (2.8 mg) half the dose being developed for PTSD – did not separate from placebo on primary endpoint, average pain improvement (responder analysis)
- Retrospective analysis showed average pain improvement (secondary endpoint) after 12 weeks of treatment showed statistical significance (P< 0.05, MMRM)
- Low dose TNX-102 SL (2.8 mg) showed an improvement in sleep quality in Phase 2 and Phase 3 FM trials
- Efficacy of TNX-102 SL 5.6 mg in FM can be studied in a potential pivotal study to support product registration

Agitation in Alzheimer's Disease

- Fast Track designation granted July 2018
- Phase 2/ potential pivotal efficacy study protocol received FDA comments in October 2018





Agitation is one of the most distressing and debilitating of the behavioral complications of Alzheimer's disease

Includes emotional lability, restlessness, irritability and aggression¹

Link between disturbed sleep and agitation in Alzheimer's¹⁻³

Agitation is commonly diurnal ("sundowning")

Prevalence

 Agitation is likely to affect more than half of the 5.3 million Americans who currently suffer from moderate to severe Alzheimer's disease, and this number is expected to nearly triple by 2050⁴

¹Rose, K.et al. (2015). American Journal of Alzheimer's Disease & Other Dementias, 30:78
 ²Shih, Y. H., et al. (2017). Journal of the American Medical Directors Association, 18, 396.
 ³Canevelli, M., et al. (2016). Frontiers in medicine, 3.
 ⁴The Alzheimer's Association, 2017 Alzheimer's Disease Facts and Figures: <u>https://www.alz.org/facts/</u>
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Consequences of Agitation in Alzheimer's Disease

57

Outcomes

 Agitation is associated with significant poor outcomes for Alzheimer's patients and challenges for their caregivers

Common reason for institutionalization

 Development of agitation, or its worsening, is one of the most common reasons for patients having to transition from lower- to higher levels of care (nursing homes and other long-term care settings)¹

Cost

 The presence of agitation nearly doubles the cost of caring for patients with Alzheimer's disease, and agitation is estimated to account for more than 12% of the healthcare and societal cost of Alzheimer's disease, which is currently estimated to be \$256 Billion for the year 2017 in the United States¹

¹The Alzheimer's Association, 2017 Alzheimer's Disease Facts and Figures: <u>https://www.alz.org/facts/</u>





FDA designated Fast Track development program

Significant unmet need

• No FDA approved drugs for the treatment of agitation in Alzheimer's

Mechanism of improving sleep quality

· Sleep disturbance is a significant and common symptoms in Alzheimer's

Pharmacological advantages outweigh potential concerns of using TNX-102 SL in treating agitation in Alzheimer's disease

Blocks 3 receptors, not just one (e.g., 5-HT_{2A})

TNX-102 SL for Agitation in Alzheimer's – Regulatory Status and Registration Strategy



FDA confirmed no additional study was needed prior to IND submission

 Pre-IND meeting established open dialogue with the FDA on pivotal clinical study design and efficacy endpoints to support product registration

Proposed Phase 2 IND study can potentially serve as a pivotal efficacy study to support NDA approval

FDA comments on final protocol received October 2018

Registration Strategy of TNX-102 SL for agitation in Alzheimer's disease

 Efficacy Supplement (sNDA¹) may be leveraged from the PTSD/FM development program and supported by Initial NDA approval for PTSD/FM

Supplemental New Drug Application

5 TNX-102 SL Potentially Addresses Some of the Challenges in Treating Agitation in Alzheimer's



Sublingual route of administration (no swallowing)

· Swallowing can be an issue for a significant number of Alzheimer's patients

Low dose taken daily at bedtime

 Potentially minimize daytime anticholinergic side effects → improved tolerability and patient compliance

Role of sleep in clearing debris from the brain

 Animal studies have shown debris clearance from the brain during sleep including toxic proteins associated with Alzheimer's progression¹

¹T Xie L, et al. Science. (2013);342(6156):373



Scientific Rationale for Developing TNX-102 SL for Agitation in Alzheimer's Disease

61

Connection between Sleep Disturbance and Agitation

- Agitation in Alzheimer's Disease is associated with sleep disturbance^{1,2}
- Evidence that improving sleep could improve agitation³

Supported by Potential Mechanism of Action

- TNX-102 is a multifunctional agent including antagonism of 5-HT_{2A}, α₁-adrenergic and histamine H₁ receptors
- Certain 5-HT_{2A} antagonists have shown clinical efficacy against agitation in dementia including trazodone^{4,5}, and mirtazapine⁶
- The α₁-adrenergic antagonist prazosin has shown efficacy in the treatment of agitation in dementia⁷
- The histamine H₁ antagonist hydroxyzine had historical use in treating agitation in dementia⁸

¹Bachmen, D. and Rabins, P. <u>Annu Rev Med.</u> 2006;57:499. ²Rose, K et al. <u>Am J Alzheimers Dis Other Demen.</u> 2015 30(1):78. ³Figueiro MG Sleep Med. 2014 15(12):1554-64. ⁴Lebert F. et al. <u>Dement Geriatr Cogn Disord.</u> 2004;17(4):355. ⁵Sulzer DL et al.<u>Am J Geriatr Psychiatry.</u> 1997 5(1):60. ⁴Cakir S. et el., <u>Keuropsychiatr Dis Treat.</u> 2008 4(5):963. ⁵Wang, LY et al., <u>Am J Geriatr Psychiatry.</u> 2019 17(9):744 ⁸Settel E. Am <u>Pract Dig Treat.</u> 1957 8(10):1584. ⁽²⁾ 2011

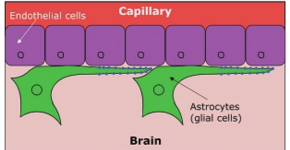
Protective Barriers in the Central and Peripheral Nervous Systems



Glial cells are cells that reside in the central nervous system and can provide protective barriers between the central and peripheral nervous systems^{1,2}

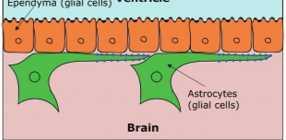
Blood-Brain Barrier:

supplies nutrients to the brain and filters toxins¹



Cerebrospinal Fluid (CSF)–Brain Barrier/Glymphatic System: extracts toxins from the brain²

Ependyma (glial cells) Ventricle



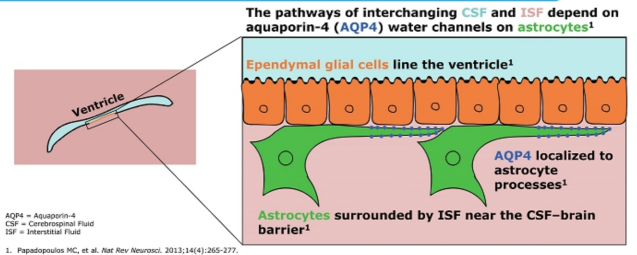
1. Ballabh P, et al. Neurobiol Dis. 2004;16(1):1-13.

Jessen NA, et al. Neurochem Res. 2015;40(12):2583-2599.
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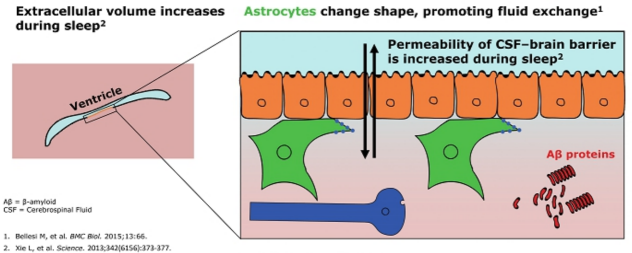


During Wakefulness, Proteins Linked to Neuronal Death and Neurodegeneration Accumulate in the Brain's Extracellular Space

63



During Sleep, the CSF–Brain Barrier Is More Permeable, Allowing Debris to Clear



64

Sleep–Wake Cycles Alter Permeability of the CSF– Brain Barrier

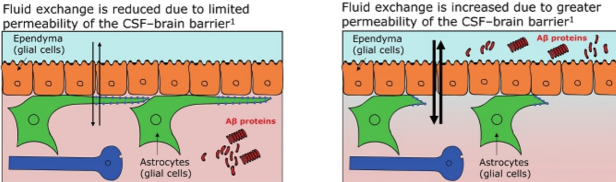
Fluid exchange at the CSF-brain barrier allows for clearance of toxic proteins called β-amyloids (Aβ).¹ Glial cells in the brain work to facilitate this fluid exchange.² Sleep-wake cycles alter glial cell morphology, which may affect fluid exchange at the CSF-brain barrier.³

Sleep:

65

Wakefulness:

Fluid exchange is reduced due to limited permeability of the CSF-brain barrier1



1. Xie L, et al. Science. 2013;342(6156):373-377. 2. Papadopoulos MC, et al. Nat Rev Neurosci. 2013;14(4):265-277. 3. Bellesi M, et al. BMC Biol. 2015;13:66. @ 2019



Agitation in Alzheimer's – Competitive Landscape of Select Drugs in Development

66

Competitive landscape

5HT_{2A} Antagonists/inverse agonists

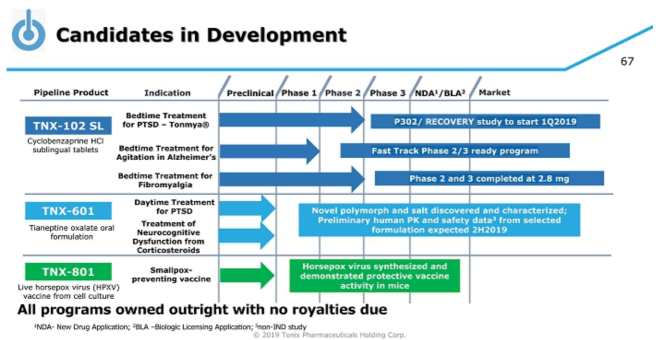
Nelotanserin (Axovant)

- Atypical Antipsychotics (also have 5HT_{2A} antagonism)
 - Rexulti® brexpiprazole (Otsuka/Lundbeck)
 - Lumateperone (Intra-Cellular)
- · Dextromethorphans believed to act as SSRI, glutamate/NMDA and sigma-1 receptor modulators
 - · Deudextromethorphan (Avanir/Otsuka) deuterated version of Nuedexta®
 - · Dextromethorphan/bupropion (Axsome Therapeutics)

TNX-102 SL uniquely designed for bedtime dosing and transmucosal absorption

- Maximize drug exposure during sleep → improving sleep quality
- Other 5-HT_{2A} antagonists not designed for bedtime sublingual dosing

NDA approval can rely on reference listed drug (AMRIX) safety information



OVERTIMATE OT A Set UP: A Potential Clinical Candidate for PTSD

Pre-IND Candidate	 Targeted as a 1st line monotherapy for PTSD: oral formulation for daytime dosing Leverages expertise in PTSD (clinical and regulatory experience, market analysis, etc.) Mechanism of Action (MOA) is different from TNX-102 SL Tianeptine sodium (amorphous) has been approved in EU, Russia, Asia and Latin America for depression since 1987 with established post-marketing experience Identified new oxalate salt polymorph with improved pharmaceutical properties ideal for reformulation Preliminary human pharmacokinetic and safety data (non-IND study) from selected formulation expected in second half 2019
Targeting a	 Issued patent on steroid-induced cognitive impairment and memory loss issues
Condition with	Clinical evidence for PTSD
Significant	 Several studies have shown tianeptine to be active in the treatment of PTSD¹⁻⁴
Unmet Need	
	Behav (hysiol. 2008 Jan;38(1):55-61. pMID: 18097761 m 5 S Korsakova. 2005;156(11):24-0. PMID: 16329631 [Russian]

68

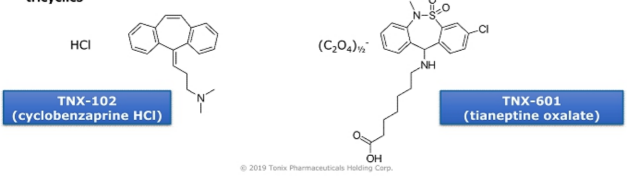




Cyclobenzaprine and tianeptine share structural similarities with classic tricyclic antidepressants (TCAs) and to each other, but each has unique pharmacological properties

· Tianeptine has a 3-chlorodibenzothiazepine nucleus with an aminoheptanoic side chain

Tianeptine leverages Tonix's expertise in the pharmacology and development of tricyclics $$\searrow_0^0$$





TNX-801 (Synthesized Live Horsepox Virus): A Smallpox-Preventing Vaccine Candidate

	Potential improvement over current biodefense tools against smallpox ✓ Leverages Tonix's government affairs effort ✓ Collaboration with Professor David Evans and Dr. Ryan Noyce at University of Alberta ✓ Demonstrated protective vaccine activity in mice
Pre-IND Stage	✓ Patent application on novel vaccine submitted Regulatory strategy
	 We intend to meet with FDA to discuss the most efficient and appropriate investigational plan to support the licensure, either:
	✓ Application of the "Animal Rule", or
	✓ Conducting an active comparator study using ACAM2000
	 Good Manufacturing Practice (GMP) viral production process in development
Tornating a	Material threat medical countermeasure under 21st Century Cures Act
Targeting a Potential Public	 Qualifies for Priority Review Voucher (PRV) upon licensure*
Health Issue	\checkmark PRVs have no expiration date, are transferrable and have sold for ~\$125 M

70

*BLA/NDA priority 6-month review is expected.

TNX-801 (Synthesized Live Horsepox Virus): A Smallpox-Preventing Vaccine Candidate



Synthesis¹ from sequence of a 1976 Mongolian isolate² In mice, TNX-801 behaved like attenuated vaccinia virus

· Vaccinia is the term used to classify the live poxviruses that are used as smallpox vaccines, including ACAM2000, which is the latest smallpox vaccine licensed in the U.S.

How is HPXV related to modern vaccines?

- Multiple sources³⁻⁵ indicate that the smallpox vaccine discovered by Dr. Edward Jenner in the early 19th century was either HPXV or a very similar virus and that vaccinia vaccines are derived from this ancestral strain
- A 1902 U.S. smallpox vaccine was found to be highly similar (99.7% similarity in core genome⁶) to HPXV sequence from the 1976 Mongolian isolate
- Horsepox is now believed to be extinct⁵

¹ Noyce, RS, Lederman S, Evans DH. PLoS ONE. 2018; 13(1): e0188453 https://doi.org/10.1371/journal.pone.0188453

Noyce, KS, Ledeman S, Evans DH. PLoS ONE: 2016; 13(1): 60160453 https://doi.org/10.13/14601616066
 ² Tulman et al., Journal of Virology, 2006; 80(18): 9244-9258
 ³ Qin et al., Journal of Virology, 2011; 85(24):13049-13060
 ⁴ Medaglia et al., Journal of Virology, 2015; 89(23):11909-11925
 ⁶ Esparza J, Veterinary Record, 2013; 173: 272-273
 ⁸ Schrick, L. et al., N Engl J Med 2017; 377:1491-1492, http://www.neim.org/doi/full/10.1056/NEJMc1707600

The Currently Licensed Smallpox Vaccine ACAM2000 is a Live Vaccinia Virus (VACV) Vaccine

ACAM2000 is sold to the U.S. Strategic National Stockpiles¹

- Sold by Emergent BioSolutions
- Sanofi divested ACAM2000 to Emergent BioSolutions in 2017 for \$97.5 M upfront plus milestones
- ACAM2000 was developed by Acambis which was acquired by Sanofi in 2008 for \$513 M

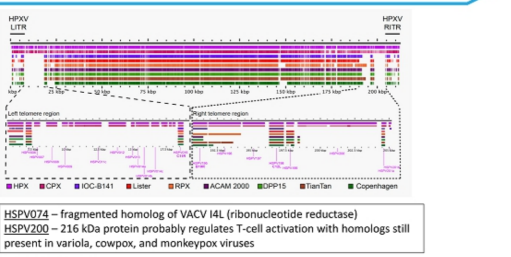
Vaccinia (VACV) strains have demonstrated potential for zoonotic infections and re-infection of humans²⁻⁵

 No known evidence for zoonosis of ACAM2000, but it has not been widely administered

Modern VACV smallpox vaccines are associated with cardiotoxicity⁶

¹Nalca, A et al. Drug design, development and Therapy. (2010) 4:71-79
 ²Medaglia MLG, et al. J Virol. (2015) 89:11909 –11925. doi:10.1128/JVI.01833-15.
 ³Trindade,GS. et al. Clinical Infectious Diseases. (2009) 48:e37-40
 ⁴Leite,JA, et al. Emerging Infectious Diseases. (2005) www.cdc.gov/eid • Vol. 11, No. 12
 ⁵Medaglia MLG, et al. Emerging Infectious Diseases (2009) www.cdc.gov/eid • Vol. 15, No. 7
 ⁶Engler RJM et al., PloS ONE (2015) 10(3): e0118283. doi:10.1371/journal.pone.0118283



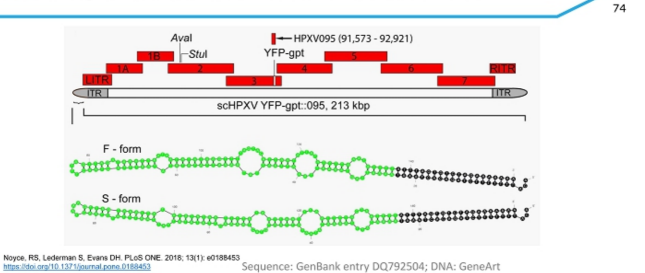


73

Evans, D. U. of Alberta (2018) with permission

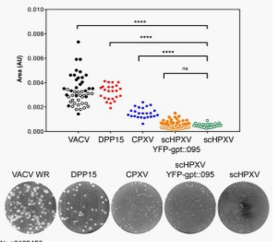


Genome Assembly (212 kbp) by Synthesis of Fragments and Construction of Telomeres





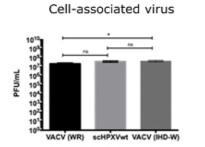
HPXV Produces Small Plaques that are More Like Cowpox Than Vaccinia (VACV)



75

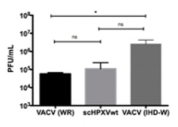
Noyce, RS, Lederman S, Evans DH. PLoS ONE. 2018; 13(1): e0188453 https://doi.org/10.1371/journal.pone.0188453 © 2019 Tonix Pharmaceuticals Holding Corp.





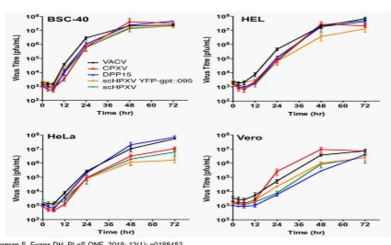
Virus in the media

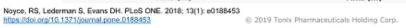
76



Noyce, RS, Lederman S, Evans DH, PLoS ONE, 2018; 13(1): e0188453 https://doi.org/10.1371/journal.pone.0188453

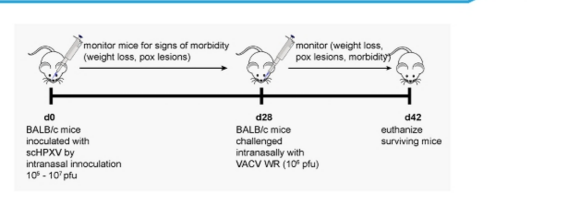








Testing Vaccine Protective Activity of HPXV in Mice Model

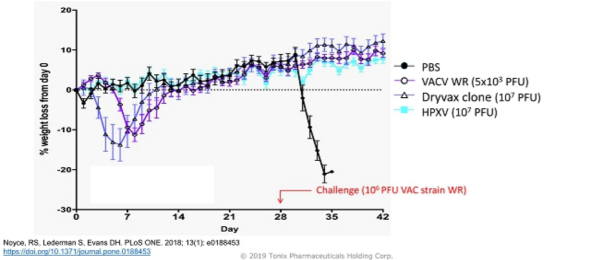


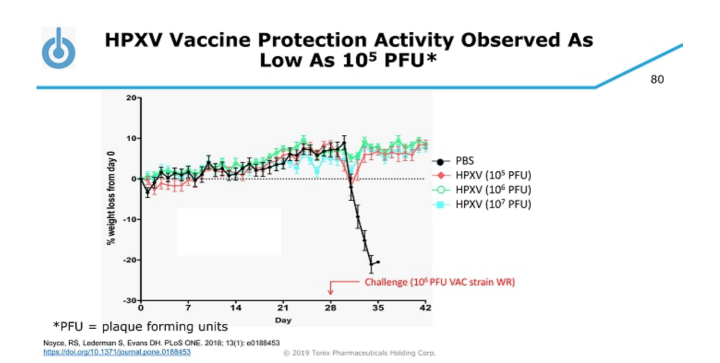
78

Noyce, RS, Lederman S, Evans DH. PLoS ONE. 2018; 13(1): e0188453 https://doi.org/10.1371/journal.pone.0188453

Biological Properties of HPXV: Less Virulent than a Dryvax Clone, but Produces Protective Immunity

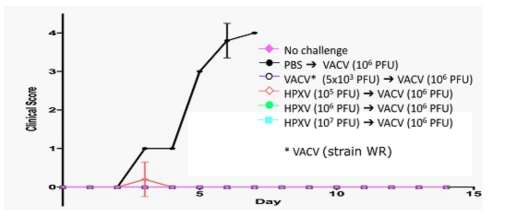
79







No Overt Clinical Sign Observed in HPXV Vaccinated Mice After VACV Challenge



Noyce, RS, Lederman S, Evans DH. PLoS ONE. 2018; 13(1): e0188453 https://doi.org/10.1371/journal.pone.0188453

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81

BALENCE OF TOX-801 – May Have an Improved Safety Profile as a Smallpox Preventing Vaccine Horsepox is caused by HPXV and is characterized by mouth and skin eruptions

HXPV isolate from the 1976 outbreak later sequenced

Modern smallpox vaccines are associated with cardiotoxicity¹

HPXV has potential for slower proliferation leading to possibly decreased toxicity²

¹ Engler RJM et al., PloS ONE 10(3): e0118283. doi:10.1371/journal.pone.0118283 (2015) ² Noyce, RS, Lederman S, Evans DH. PLoS ONE. 2018; 13(1): e0188453 <u>https://doi.org/10.1371/journal.pone.0188453</u>



An Improved Smallpox-Preventing Vaccine is Important and Necessary for a Potential Public Health Issue

Smallpox was eradicated as a result of global public health campaigns 83

No cases of naturally-occurring smallpox have been reported since 1977

Accidental or intentional transmission of smallpox does not require a natural reservoir

Stockpiles of smallpox-preventing vaccines are currently maintained and refreshed in case of need



Ongoing vaccination of U.S. troops

Troops in the Global Response Force

Threat of smallpox re-introduction

Strategic National Stockpile & public health policy

Re-emergence of monkey pox¹

· Believed to resurgent because of vaccinia-naïve populations in Africa

84

Multiple U.S. military operations ongoing in Africa

¹Nda- Isaiah, J. Nigeria: Monkey Pox Scourge Spreads to Seven States. All Africa. 12 OCTOBER 2017, <u>HTTP://ALLAFRICA.COM/STORIES/201710120177.HTML</u> © 2019 Tonix Pharmaceuticals Holding Corp.



21st Century Cures Act (2016), Section 3086

· Encouraging treatments for agents that present a national security threat

85

Medical countermeasures are drugs, biologics (vaccines) or devices intended to treat:

- Biological, chemical, radiological, or nuclear agents that present a national security threat
- Public health issues stemming from a naturally occurring emerging disease or a natural disaster

New Priority Review Voucher program for "Material Threat Medical Countermeasures"

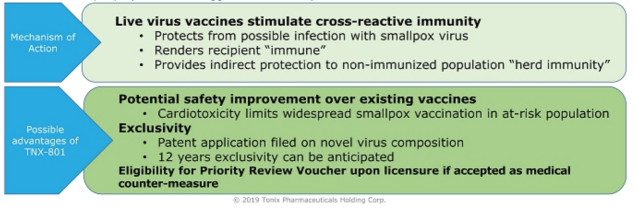
Priority Review Voucher may be transferred or sold
 © 2019 Tonix Pharmaceuticals Holding Corp.

6

TNX-801 (Synthesized Live Horsepox Virus): A Smallpox-Preventing Vaccine Candidate

TNX-801 (HPVX)

- Synthesized live horsepox virus
- · Shares structural characteristics with vaccinia-based smallpox vaccines
- · Unique properties that suggest lower toxicity



86



Given that smallpox is eradicated the only evidence of effectiveness for modern vaccines is from historical use when smallpox was endemic

87

· Stimulates interest in the evolution of vaccinia

Vaccinia stocks around the world diverged from Jenner's 1798 vaccine

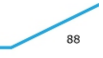
Evolutionary argument that common progenitor was horsepox or a similar virus

U.S. vaccine from 1902 was found to be 99.7% similar to horsepox in core viral sequence¹

- · Strong evidence linking a horsepox-like virus as progenitor to modern vaccinia
- Effectiveness of older vaccines support belief that HPXV will be protective against smallpox

¹Schrick, L. et al (2017) An Early American Smallpox Vaccine Based on Horsepox N Engl J Med 2017; 377:1491 © 2019 Tonix Pharmaceuticals Holding Corp.





Single clone picked from "swarm" of Dryvax^{®1}

• Some rationale for selection²

Growth in serum free Vero cells

 Eliminates risk of Bovine Spongiform Encephalopathy (BSE)/prion contamination – safety concerns in Wyeth's Dryvax (grown in calf lymph)

In 2000, the evolutionary connection between vaccinia and horsepox was not understood

Tulman's sequence of horsepox was published in 2006³

¹US licensed smallpox preventing vaccine - ACAM2000 is currently marketed, Dryvax has been withdrawn from marketing
 ²Monath, TP et al. Int. J. of Inf. Dis. (2004) 852:S31
 ³Tulman, ER. Genome of Horsepox Virus J. Virol. (2006) 80(18) 9244
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Rationale for Developing a Potentially Improved New Smallpox Vaccine



Toxicity concern of modern vaccinia (VACV) vaccines limit wildly administration

- · Not recommended for use, even in first responders
- · U.S. soldiers in the Global Response Force are immunized

Modern VACV vaccination safety studied in 1081 VACV (Dryvax [62.5%] and ACAM2000 [37.5%]) vaccinees¹

- New onset chest pain, dyspnea and/or palpitations 10.6% of VACV-vaccinees and 2.6% of control immunized (TIV)²
- Clinical: 4 probable myo- and 1 suspected peri-carditis (5 cases out of 1081 VACV vaccinees – 0.5%)
- Cardiac specific troponin T (cTnT) elevation in 31 VACV vaccinees (3%)

¹Engler RJM,, et al. (2015) A Prospective Study of the Incidence of Myocarditis/Pericarditis and New Onset Cardiac Symptoms following Smallpox and Influenza Vaccination. PLoS ONE 10(3) ²TIV = trivalent influenza vaccine - control vaccinees © 2019 Tonix Pharmaceuticals Holding Corp.

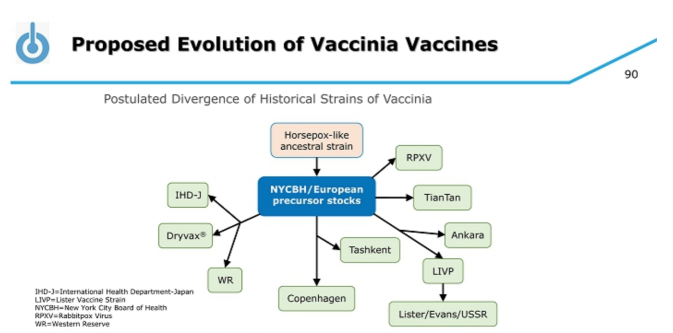
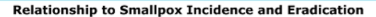
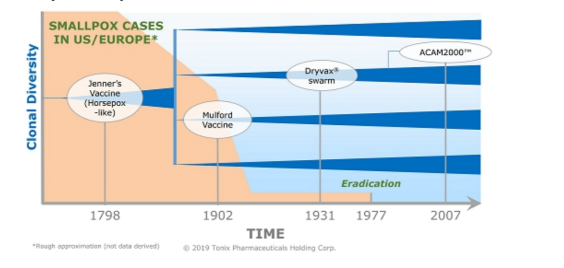


Figure Adapted from Qin et al. Journal of Virology. 2015;89(3):1809-1824. © 2019 Tonix Pharmaceuticals Holding Corp.









What's the Evidence of Effectiveness of Smallpox Vaccines for Preventing Smallpox?

Theoretical effectiveness of modern vaccinia vaccines are based on extrapolation from older vaccines

· Newer/modern vaccines were not widely used when smallpox was endemic

92

MVA (Modified Virus Ankara) which has large deletions also produces different T cell responses

- In non-human primates, MVA is less effective than ACAM2000 in protecting against monkeypox¹
- MVA has fewer epitopes, and elicits different responses to existing epitopes²
 - MVA effectiveness argument is based on the immune response to intracellular mature virus (IMV)
 - Immunity to the other form of virus, extracellular enveloped virus (EEV), is weak because the immunodominant B5 gene is heavily mutated and deleted in MVA

¹Golden JW, et al. (2012). PLoS ONE 7(7): e42353. doi:10.1371/journal.pone.0042353 ²Tscharke, DC et al., J. Exp. Med. 2005 201(1):95 (2019 Tonix Pharmaceuticals Holding Corp.

Possible Smallpox Prevention and Treatment Strategies

Preventing Vaccine

· Jenner's vaccine, HPXV (upon licensure), Vaccinia

Post-exposure vaccination¹

Jenner's vaccine

Priming of the immune system

Imvamune[®] (MVA) and DNA vaccines²

Pharmacotherapy for infected or exposed individuals

Arestvyr®/TPOXX® (tecovirimat, formerly ST-246)

Treatment of disseminated viremia in immunocompromised³

Arestvyr[®]/TPOXX[®], Brincidofovir and vaccinia immune globulin

¹Described by Jenner as one of his major discoveries

²Hooper, JW et al. Smallpox DNA Vaccine Protects Nonhuman Primates Against Lethal Monkeypox. J. Virol. 2004. 78 (9) 4433
³Lederman, ER et al, Progressive Vaccinia: Case Description and Laboratory-Guided Therapy With Vaccinia Immune Globulin, ST-246, and CMX001 JID 2012. 206:1372

93



Viral Replication Proficiency is Critical to Human Immunogenicity but May Compromise Safety

Pox vaccines with low or no replication appear safer than vaccines replicate fast in human cells

 Canarypox and Imvamune[®] (Modified Virus Ankara/MVA) appear to have good tolerability

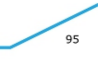
94

- · Relatively safe in immunocompromised hosts
- Rapidly replicating modern vaccinia vaccines (Dryvax® and ACAM2000®) are associated with myocarditis

Replication correlates positively with immunogenicity

- · Jenner's vaccine and modern vaccinia engender strong immunity
- Canarypox and MVA appear to be weak immunogens, suitable for priming of the immune system in healthy human being and potentially safe enough to use in immunocompromised people





TNX-801 (HPXV) is expected to have similar scalability for mass production as ACAM2000

- TNX-801 grows well in cell lines immunity is expected after single administration (immunization)
- · Only a small dose (replicating live virus) is required for immunization

MVA is hard to scale up for commercial production

- Requires high dose to engender an immune response (non-replicating virus)
- Cumbersome immunization schedule two doses, 4 weeks apart, are used typically to prime the immune system (slow growth)

Antivirals

- · Relatively expensive to manufacture requires repeated dosing
- · May provide logistical challenges to at risk population over the at risk period



Rationale for Developing a Potentially Improved New Smallpox Vaccine Based on Jenner's Vaccine

96

Vaccination protects against smallpox – both individuals and populations at risk

Use of Jenner's vaccine resulted in eradication of smallpox

Vaccination can protect AFTER smallpox infection

Vaccinia can be administered 1-3 days after infection

Vaccination indirectly protects non-immunized people in a population

"Wetting the forest" or "herd immunity"

Vaccination can be cost effective with safe/low-risk vaccines

 Replication-efficient live virus vaccines can be manufactured and administered for broader use

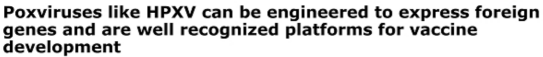
"The Time is Right"

New synthetic biology technology and new understanding of vaccinia evolution provide an opportunity for a potentially safer vaccine using HPXV



Potential for Use of HPXV as a Vector for Vaccines to Infectious Disease or Cancer

97



- Large packaging capacity for exogenous DNA inserts (i.e. encoding antigens)
- · Precise virus-specific control of exogenous gene insert expression
- · Lack of persistence or genomic integration in the host
- · Strong immunogenicity as a vaccine
- · Ability to rapidly generate vector/insert constructs
- · Readily manufacture at scale
- · Live, replicating vaccine direct antigen presentation

Potential advantages of HPXV- strong immunogenicity with good tolerability



NASDAQ: TNXP

Cash and cash equivalents, September 30, 2018	\$14.8 million
Net proceeds from equity offerings in 4Q2018	\$17.3 million
Common Stock outstanding post December 2018 underwritten equity offering (as of March 1, 2019)	4.7 million
Pro Forma Common Stock outstanding post December 2018 underwritten equity offering ¹ (as of March 1, 2019)	6.1 million

98

¹ Pro forma to include the remaining 1.4 million shares of Common Stock, not yet converted as of March 1, 2019, issuable upon conversion of the Series A Convertible Preferred Stock, as per terms of the December 2018 underwritten offering.



Management Team





Board of Directors



Donald Landry, MD, PhD Chair of Medicine, Columbia University Adeoye "Oye" Olukotun, MD Squibb, BMS, Mallinckrodt, Esperion	
James Treco First Chicago, Salomon Brothers/Citigroup	

b Milestones – Recently Completed and Upcoming

ď	July 2018	Completed P301/HONOR study interim analysis - result did not support study continuation but strengthened new Phase 3 study
ন্দ্র	August 2018	Presentation of P301/HONOR study results at Military Health System Scientific Symposium
র্জ	October 2018	Met with FDA and received preliminary agreement on the design of new Phase 3 study of Tonmya for PTSD (P302/RECOVERY study)
⊻	November 2018	Received FDA minutes confirming agreement on the design of P302/RECOVERY study
	First Quarter 2019	P302/RECOVERY study to be initiated
	First Quarter 2019	FM FDA Clinical Guidance meeting
	Second Half 2019	Preliminary human pharmacokinetic and safety data (non-IND study) from selected TNX-601 (tianeptine oxalate) formulation expected
	First Half 2020	Topline data from P302/RECOVERY study expected

101



Summary



Phase 3 development of new bedtime treatment for PTSD, including militaryrelated PTSD

- · Major unmet need; ~12 million Americans annually
- · Benefited from FDA 505(b)(2) NDA approval requirement

Complimentary day-time PTSD treatment in development

 Leverages development expertise in PTSD, i.e., regulatory, trial recruitment and execution

New indication in development for agitation in Alzheimer's Disease

- · Unmet medical need, no approved drug available
- Fast Track Phase 2/3 ready program

Fibromyalgia bedtime treatment in development

· IND ready to support Phase 3 potential pivotal efficacy study

Innovative vaccine in development to prevent Smallpox

- · Opportunity to supply stockpiling requirement; short development path
- · Studies in mice suggest improved safety profile







Exhibit 99.02





March 2019

Version P0160 3-5-19 (Doc 0438)



Cautionary Note on Forward-Looking Statements

2

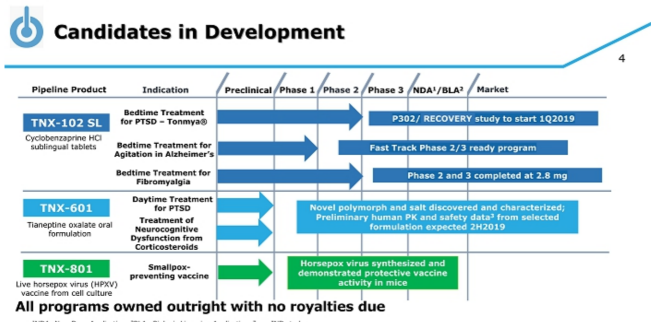
Certain statements in this presentation regarding strategic plans, expectations and objectives for future operations or results are "forward-looking statements" as defined by the Private Securities Litigation Reform Act of 1995. These statements may be identified by the use of forward-looking words such 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, substantial competition; 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 risks related to failure to obtain U.S. Food and Drug Administration clearances or approvals and noncompliance with its regulations. 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, 2017, as filed with the Securities and Exchange Commission (the "SEC") on March 9, 2018, and periodic 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.



Posttraumatic Stress Disorder (PTSD) - Lead program; new bedtime Cyclobenzaprine Sublingual Tablets treatment - Tonmya®1 **INX-102 SI** P302/RECOVERY Phase 3 clinical study with Week 4 primary endpoint to initiate in 1Q2019 · Results from 2 efficacy studies improve the new Phase 3 study design New Phase P302/RECOVERY study design features accepted by the FDA² Agitation in Alzheimer's disease (AAD) IND³ ready to support Phase 2 potential pivotal efficacy study Fibromyalgia Syndrome (FM) IND³ ready to support Phase 3 potential pivotal efficacy study TNX-601⁴ – Daytime PTSD treatment and treatment of neurocognitive Pipeline dysfunction from corticosteroids Pre-IND candidate; nonclinical development ongoing TNX-801⁵ - Smallpox-preventing vaccine candidate Efficacy demonstrated in mouse model ¹ Tonmya has been conditionally accepted by the U.S. FDA as the proposed trade name for TNX-102 SL (cyclobenzaprine HCI sublingual tablets) for the treatment of PTSD. TNX-102 SL is an investigational new drug and has not been approved for any indication. ² FDA Meeting Minutes (November 26, 2018) ³ IND- Investigational New Drug Application ⁴ Timeetine evaluate

3

⁴ Tianeptine oxalate ⁵ Synthesized live horsepox virus



¹NDA- New Drug Application; ²BLA -Biologic Licensing Application; ³non-IND study (© 2019 Tonix Pharmaceuticals Holding Corp.

Tonmya: a Potential Bedtime Treatment for PTSD

First investigational new drug to show treatment effect in military-related PTSD in two potential pivotal efficacy studies

Phase 2 study (P201/AtEase) showed Tonmya 5.6 mg had a strong signal of treatment effect at Week 12 as measured by CAPS-51

5

- Phase 3 study (P301/HONOR) provided evidence of effectiveness as early as 4 weeks after treatment but diminished over time due to high placebo response
 - Retrospective analysis showed persistent effectiveness at Week 12 in subgroup with trauma ≤9 years from screening
- · Both studies can be used as supportive evidence of efficacy and safety for Tonmya NDA submission
- No serious or unexpected adverse events related to Tonmya were reported

FDA feedback and acceptance on new Phase 3 study (P302/RECOVERY) received in November² Patent protection through 2034 in U.S.³

Composition of matter patent for transmucosal delivery of cyclobenzaprine

Novel mechanism targets sleep quality

Memory processing during sleep is important to recovery from PTSD

CAPS-5 = Clinician-Administered PTSD Scale for DSM-5
 FDA Meeting Minutes, November 26, 2018; ³U.S. Patent No. 9,636,408 for eutectic proprietary ProtecticTM formulation





Active ingredient is cyclobenzaprine, which is structurally related to tricyclic antidepressants

- Cyclobenzaprine interacts with receptors that regulate sleep quality: $5-HT_{2A_{1}} \alpha_{1}$ -adrenergic and histamine H₁ receptors
- Cyclobenzaprine does <u>NOT</u> interact with the same receptors as traditional hypnotic sleep drugs, benzodiazepines or nonbenzodiazepines that are associated with retrograde amnesia
- Cyclobenzaprine-containing product was approved 40 years ago and current labeling (May 2018) indicates no abuse or dependence concern

Tonmya NDA can be filed without drug abuse and dependency assessment studies

· Discussed at March 9, 2017 meeting with the FDA



TNX-102 SL Intellectual Property – U.S. Protection until 2034

Composition of matter (eutectic) : Protection expected to 2034

- United States Patent and Trademark Office (USPTO) issued U.S. Patent No. 9,636,408 in May 2017 U.S. Patent No. 9,956,188 in May 2018 and U.S. Patent No. 10,117,936 in November 2018
- Japanese Patent Office (JPO) issued Japanese Patent No. 6310542 in March 2018
- New Zealand Intellectual Property Office (NZIPO) issued New Zealand Patent No. 631152 in May 2017

7

37 patent applications pending (2 allowed (US and South Africa))

Pharmacokinetics (PK) : Protection expected to 2033

- JPO issued Japanese Patent No. 6259452 in December 2017
- NZIPO issued New Zealand Patent No. 631144 in March 2017
- Taiwanese Intellectual Property Office issued Taiwanese Patent No. I590820 in July 2017
- · 21 patent applications pending (1 allowed (Australia))

Method of use for active ingredient cyclobenzaprine : Protection expected to 2030

- USPTO issued U.S. Patent 9,918,948 in March 2018
- European Patent Office issued European Patent No. 2 501 234B1 in September 2017 (validated in 38 countries). Opposition filed in June 2018
- · 2 patent applications pending



TNX-102 SL: Sublingual Formulation is Designed for Bedtime Administration



8

- · Innovation by design with patent protected CBP/mannitol eutectic
- Rapid systemic exposure
- · Increases bioavailability during sleep
- Avoids first-pass metabolism
- · Lowers exposure to long-lived active major metabolite, norcyclobenzaprine (norCBP)

CBP undergoes extensive first-pass hepatic metabolism when orally ingested

- Active major metabolite, norCBP¹
 - Long half-life (~72 hours)
 - Less selective for target receptors (5-HT_{2A}, α₁-adrenergic, histamine H₁)
 - More selective for norepinephrine transporter and muscarinic M₁

TNX-102 SL 505(b)(2) NDA approval can rely on the safety of the reference listed drug (AMRIX[®])²

¹ Daugherty et al., Abstract 728, Society of Biological Psychiatry 70th Annual Scientific Convention, May 14-16, 2015, Toronto Ontario, Canada ² FDA Minutes (November 26, 2018) © 2019 Tonix Pharmaceuticals Holding Corp.



Tonmya: Hypothesized Novel Mechanism Targets Sleep Quality for Recovery from PTSD

PTSD is a disorder of recovery

- · Most people exposed to extreme trauma recover over a few weeks
- In PTSD, recovery process impeded due to insufficient sleep-dependent memory processing^{1,2}

9

Memory processing is essential to recovery

 Vulnerability to memory intrusions and trauma triggers remains if no consolidation of new learning (extinction)

Tonmya targets sleep quality³

 The active ingredient in Tonmya, cyclobenzaprine, interacts with receptors that regulate sleep quality: strongly binds and potently blocks 5-HT_{2A}, α₁-adrenergic and histamine H₁ receptors, permissive to sleep-dependent recovery processes

¹Straus LD, Acheson DT, Risbrough VB, Drummond SPA. Sleep Deprivation Disrupts Recall of Conditioned Fear Extinction. Biol Psychiatry Cogn Neurosci Neuroimaging. 2017; 2(2):123-129. "Murkar ALA, De Koninck J. Consolidative mechanisms of emotional processing in REM sleep and PTSD. Sleep Med Rev. 2018; 41:173-184. "Daugherty et al., Abstract 728, Society of Biological Psychiatry 70th Annual Scientific Convention, May 14-16, 2015, Toronto Ontario, Canada



FDA-approved SSRIs, paroxetine and sertraline, are indicated as a treatment for PTSD

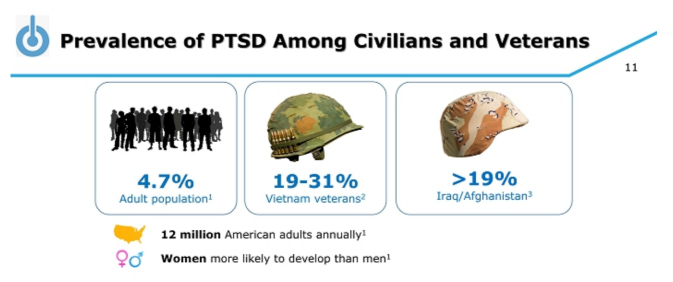
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- · Neither drug has shown efficacy in military-related PTSD
- · Majority of male PTSD patients unresponsive or intolerant to current treatments
- Side effects relating to sexual dysfunction, sleep disturbance and weight gain are commonly reported

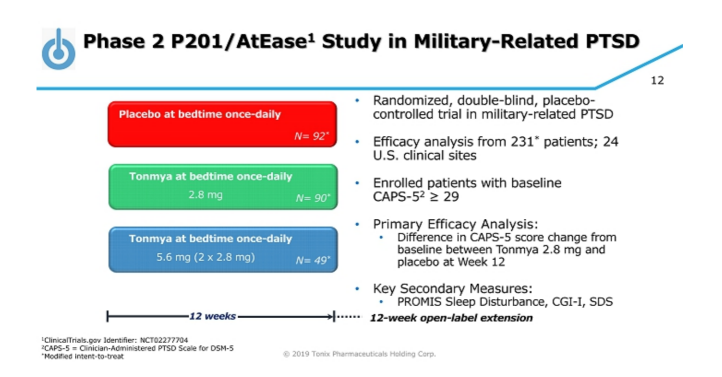
Characteristics of an ideal drug therapy that would be compatible and complementary with behavioral therapy

- Lack of retrograde amnesia (e.g., unlike off-label use of benzodiazepines and nonbenzodiazepines)
- · Lack of interference on sleep (e.g., unlike approved SSRIs)

Tonmya is being investigated in both military and civilian PTSD and is expected to be indicated as a "treatment for PTSD"



³Goldstein et al., 2016 (adjusted for 2019); ²Norris, PTSD Res Quar. 2013; ³Analysis of VA Health Care Utilization among Operation Enduring Freedom, Operation Iragi Freedom, and Operation New Dawn Veterans, from 1st Qtr FY 2002 through 2nd Qtr FY 2015, Washington, DC; Among 1.9M separated OEF/OIF/OND veterans, 1.2M have obtained VA healthcare; 685k evaluated by VA with possible mental disorder, and 379k diagnosed with PTSD.







P201 was a large adequate well-controlled Phase 2 study in militaryrelated PTSD

- Primary endpoint (Week 12 CAPS-5) did not separate from placebo for TNX-102 SL 2.8 mg
- · No safety or tolerability issue discovered
- Retrospective analyses showed TNX-102 SL 5.6 mg had a strong signal of treatment effect at Week 12 CAPS-5 (P=0.053) and CGI-I (P=0.041) scores
- Retrospective analyses suggested CAPS-5 ≥ 33 enrollment criteria for Phase 3

P301/HONOR¹ Study – Evidence of Efficacy at Week 4 Discontinued Due to High Placebo Response at Week 12

General study characteristics:

Randomized, double-blind, placebo-controlled, adaptive design, planned 550 military-related PTSD participants with baseline CAPS- $5^2 \ge 33$ in approximately 40 U.S. sites

Primary endpoint CAPS-52:

 Mean change from baseline at Week 12 (Tonmya 5.6 mg vs. placebo)

14

Tonmya once-daily at bedtime $5.6 \text{ mg} (2 \times 2.8 \text{ mg tablets}) \qquad N= 125^{+}$ Placebo once-daily at bedtime $N= 127^{*}$

– 12 weeks -

Unblinded interim analysis at 274 randomized participants (mITT* N= 252)

- Study stopped due to not meeting a pre-specified study continuation threshold at Week 12
- Participants discontinued in HONOR or 12-week open-label extension (OLE) studies can enroll in the 40-week OLE study

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¹ClinicalTrials.gov Identifier: NCT03062540 ²CAPS-5 = Clinician-Administered PTSD Scale for DSM-5 *Modified intent-to-treat

P301/HONOR Study Stopped After Interim Analysis (July 2018)

P301 was a large adequate well-controlled Phase 3 study in militaryrelated PTSD

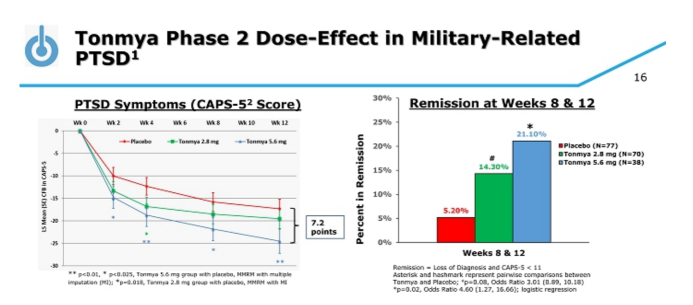
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- Separation on primary endpoint at Week 12 did not cross pre-specified study continuation threshold at Week 12 (p=0.602)
- · No safety or tolerability issue discovered
- Retrospective analyses showed Week 4 CAPS-5 (P=0.019) and CGI-I (P=0.015) scores in Tonmya group had a strong signal of treatment effect

P301 dataset is complex and rich

- Retrospective analyses presented at Military Health System Research Symposium (MHSRS) in Kissimmee, FL on August 22, 2018
- Results discussed with the FDA¹ and helped to design the new Phase 3 P302/RECOVERY study with high probability of success

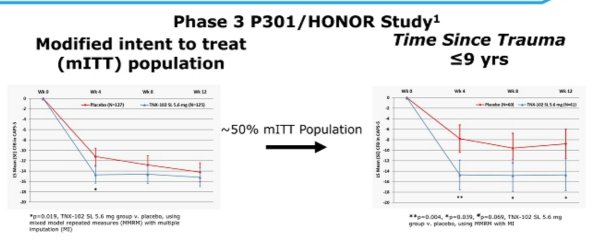
¹FDA Meeting Minutes (November 26, 2018)



³Completed Phase 2 P201/AtEase study: Retrospective analysis of Tonmya 5.6 mg on CAPS-5 ≥33 (high-moderate) subgroup. Primary analysis of P201/AtEase was on Tonmya 2.8 mg in participants with entry CAPS-5 ≥29, moderate PTSD severity. ³Clinician administered PTSD Scale for DSM-5

Primary Outcome (CAPS-5) in Phase 3 (mITT) and ≤9 Years Time Since Trauma (TST) Subgroups

17

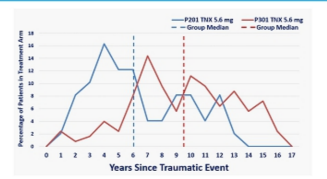


¹Phase 3 P301/HONOR study: stopped in July 2018. Separation on primary endpoint did not cross pre-specified study continuation threshold at Week 12 in the interim analysis at ~50% randomization; no safety or tolerability issues discovered. © 2019 Tonix Pharmaceuticals Holding Corp.



Retrospective Comparison of Time Since Trauma in P201/AtEase versus P301/HONOR (Tonmya 5.6 mg Groups)

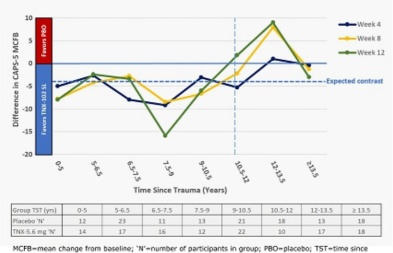
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P301 study was initiated approximately two years later than Phase 2 P201

 The median time since trauma in Phase 3 was 9.5 years compared to the median time since trauma in Phase 2 of 6.0 years for TNX-102 SL 5.6 mg treated groups

CAPS-5 Mean Change from Baseline Difference from Placebo of Tonmya 5.6 mg in TST Subgroups in P301¹



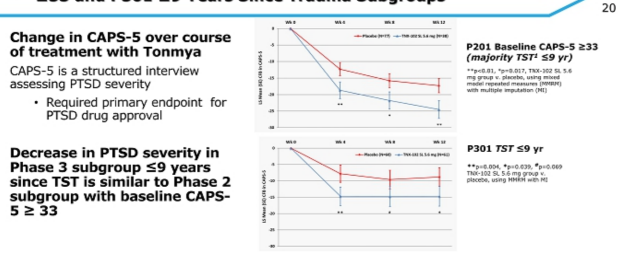
 The mITT population was divided into subgroups based on TST (1.5-2 years each as well as 0-5 years and ≥13.5 years subgroups)

19

- Graph shows the CAPS-5 differences in MCFB between TNX 5.6 mg and PBO for Weeks 4, 8, and 12 post-baseline timepoints
- and 12 post-baseline timepoints
 "Expected contrast" horizontal dashed line indicates observed effect from Phase 2 P201 study
- For TST <10.5 years groups, TNX 5.6 mg showed good separation from PBO (left side of vertical dashed 10.5 year line)
- For TST >10.5 years groups, separation of TNX 5.6 mg from PBO was either small or worked in the favor of PBO (right side of vertical dashed 10.5 year line)

¹Time Since Trauma in PTSD: Phase 3 Multi-Center, Double-Bilind, Placebo-Controlled Trial of TNX-102 SL, a Sublingual Formulation of Cyclobenzaprine, in Military-Related PTSD (Study TNX-CY-P301) Presented at CNS Summit in Boca Raton, FL November 1-4, 2018 and abstract published in Innovations in Clinical Neuroscience, November-December 2018;15(11-12, suppl):S10. https://content.equisolve.net/tonixpharma/media/1d0c405 5b2863fc74e1ef45/9ddaf42b.pdf

PTSD Treatment Response to Tonmya in Phase 2 and Phase 3 Studies: Retrospective Analyses of P201 Entry CAPS-5 ≥33 and P301 ≤9 Years Since Trauma Subgroups



¹Time since trauma; ²Majority of P201 participants were ≤9 years since trauma and ~80% of P201 participants and all of P301 participants were ≥33 CAPS-5 at baseline

6

Sustained Remission in Phase 2 and Phase 3 Studies: Retrospective Analyses of P201 Entry CAPS-5 ≥33 and P301 ≤9 Years Since Trauma Subgroups

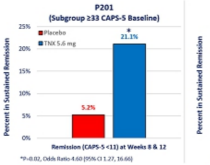
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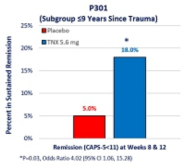
Remission is a clinical state that is essentially asymptomatic

In order to confirm remission:

 Determined rates of participants who met remission status at both Week 8 and Week 12

Rate of remission in ≤9 years since trauma group in P301 is similar to baseline CAPS-5 ≥ 33 group in P201¹





¹Majority of P201 participants were ≤ 9 years since trauma and ~80% of P201 participants and all of P301 participants were ≥ 33 CAPS-5 at baseline



Betrospective Analyses of ≤9 Years Since Trauma Subgroup on Primary and Secondary Endpoints in P301/HONOR Study

			P301 mITT			P301 ≤9 Year Subgroup				
			PBO (N=127) v. TNX-5.6 (N=125)			PBO (N=60) v. TNX-5.6 (N=61)				
			Week 4 Week 12		ek 12	Week 4		Week 12		
	Measure	Analysis	LSMD	p-value	LSMD	p-value	LSMD	p-value	LSMD	p-value
1°	CAPS-5	MMRM/MI	-3.6	0.019	-1.0	0.602	-6.9	0.004	-5.9	0.039
2°s	CGI-I	MMRM	-0.3	0.015	-0.1	0.403	-0.6	0.002	-0.5	0.021
	SDS	MMRM	-0.2	0.785	-1.6	0.101	-1.8	0.167	-4.3	0.007
	PGIC	MMRM	-0.2	0.238	-0.3	0.020	-0.4	0.045	-0.6	0.007
	PROMIS SD	MMRM	-3.1	0.015	-2.7	0.082	-4.5	0.029	-5.0	0.042
	BDI-II	MMRM	-1.1	0.330	-1.4	0.255	-5.2	0.008	-6.6	0.001

BOLDED p-values are all p<0.05; BDI-II=Beck Depression CAPS-5=Clinician-Administered PTSD Scale for DSM-5; CGI-I=Clinical Global Impression - Improvement scale; mITT=modified Intent-to-Treat sample; MNRM=mixed model repeated measures analysis; MI=multiple imputation; PGIC=Patient Global Impression of Change scale; RROMIS SD=Patient-Reported Outcome Measurement Information System Sleep Disturbance instrument (short form 8a); P80=placebo; SDS=Sheehan Disability Scale; TNX-5.6=TNX-102 SL 5.6 mg; yrs=years; 1°=primary; 2°s=secondaries

Secondary endpoints also showed strong treatment effects in ≤9 yrs TST

· Support CAPS-5 results and similar to Phase 2 P201 Study results



Adverse Events (AEs) in P201/AtEase and P301/HONOR Studies

	P201			P301	
Category of Adverse Reaction Preferred Term	Placebo (N=94)	TNX 2.8 mg (N=93)	TNX 5.6 mg (N=50)	Placebo (N=134)	TNX 5.6 mg (N=134)
Systemic Adverse Events**					
Somnolence	6.4%	11.8%	16.0%	9.0%	15.7%
Dry mouth	10.6%	4.3%	16.0%		
Headache	4.3%	5.4%	12.0%		
Insomnia	8.5%	7.5%	6.0%		
Sedation	1.1%	2.2%	12.0%		
Local Administration Site Reactions*					
Hypoaesthesia oral	2.1%	38.7%	36.0%	1.5%	37.3%
Paraesthesia oral	3.2%	16.1%	4.0%	0.7%	9.7%
Glossodynia	1.1%	3.2%	6.0%		
Product Taste Abnormal				3.0%	11.9%

23

²only adverse events (AEs) are listed that are at a rate of \geq 5% in any TNX-treated group *no values in a row for either study means the AE in the active group(s) in that study was at a rate of <5%

No serious or unexpected AEs in P201 or P301 related to Tonmya

- Systemic AEs comparable between studies and also consistent with those described in approved oral cyclobenzaprine product labeling
- Severity and incidence of oral hypoesthesia (oral numbness) are not dose related and similar in both studies
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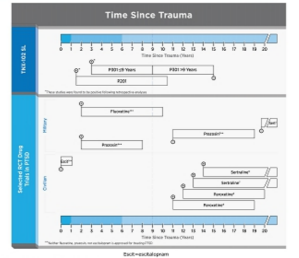
Time Since Trauma – Review of Published Studies

Published studies of prazosin suggested effects in military-PTSD prior to 9 years • Loss of treatment effect >9 years Paroxetine and sertraline studies supporting FDA approval were

conducted on PTSD > 9 years

 SSRIs have a benefit long after trauma

¹Martenyi et al. J Clin Psychiatry 2002;63:199-205.
 ²Friedman et al. J Clin Psychiatry 2007;68:711-720.
 ³Raskind et al. ACM 2010;378:507-517.
 ⁴Raskind et al. ACM J Psychiatry 2012;709:1003-1010.
 ⁵Shalev et al. ACM Gen Psychiatry 2012;69:166-176.
 ⁵Davidson et al. ACM Gen Psychiatry 2001;58:485-492.
 ³Brady et al. JAMA 2000;283:1837-1844.
 ⁴Marshall et al. ACM J Psychiatry 2001;58:1882-1988.
 ⁵Tucker et al. J Clin Psychiatry 2001;62:860-868.



24



Time Since Trauma – Remitting and Persistent Phases of PTSD

Kessler et al¹ studied remission in PTSD with and without therapy

- · Identified remitting and persistent phase of PTSD with transition at approximately 6 years post trauma
- Supported by other studies2-6 ٠

 ¹Kessler et al. Arch Gen Psychiatry 1995;52:1048-1060.

 ²Armenta et al. BMC Psychiatry 2018;18:48.

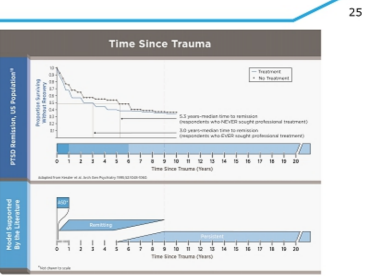
 ³Salatzer-Levy et al. PLOS OWE 2013;8:e70084.

 ³Perkoning et al. Arch Schröder 2005;162:1320-1327.

 ³Santiago et al. PLOS OWE 2013;8:e59236.

 ³Davidson & Connor. Eur Neuropsychopharmacol 2001;11(Supp3):5148-5149

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Response to Tonmya for Female Participants in P301/HONOR Study¹



Females made up only 11% of the P301/HONOR study mITT population

Difference in mean change from baseline in CAPS-5 in females between placebo (N=17) and Tonmya 5.6 mg (N=10) was:

- At 4 weeks -11.5 points
- · At 12 weeks -9.1 points

Indicates substantial separation from placebo in the small number of female participants

Predicts therapeutic response to Tonmya 5.6 mg likely in mixed civilian and military PTSD population to be studied in upcoming P302/RECOVERY trial

Civilian PTSD population tends to be about 2/3 female

¹Time Since Trauma in PTSD: Phase 3 Multi-Center, Double-Blind, Placebo-Controlled Trial of TNX-102 SL, a Sublingual Formulation of Cyclobenzaprine, in Military-Related PTSD (Study TNX-CY-P301) Presented at CNS Summit in Boca Raton, FL November 1-4, 2018 and abstract published in Innovations in Clinical Neuroscience, November-December 2018;15(11-12,suppl):S10.<u>https://content.eguisolve.net/tonixpharma/media/1d0c4055b2863fc74e1ef45f9ddaf42b.pdf</u>

Response to Tonmya for Non-Combat Traumas in P301/HONOR Study in ≤9 Years Time Since Trauma Subgroup¹

Non-combat traumas studied are similar to traumas experienced in civilian populations with PTSD

To determine the therapeutic effects of Tonmya 5.6 mg in a mixed civilian and military population, difference in MCFB in CAPS-5 was assessed in non-combat traumas in \leq 9 years TST subgroup (placebo N=14, Tonmya 5.6 mg N=10):

27

- At 4 weeks -4.8 points
- At 12 weeks -4.4 points

Non-combat traumas treated with Tonmya 5.6 mg showed clinically meaningful separation from placebo at Weeks 4 and 12, suggesting a mixed civilian and military sample within 9 years of index trauma may show a therapeutic response to Tonmya

¹Time Since Trauma in PTSD: Phase 3 Multi-Center, Double-Blind, Placebo-Controlled Trial of TNX-102 SL, a Sublingual Formulation of Cyclobenzaprine, in Military-Related PTSD (Study TNX-CY-P301) Presented at CNS Summit in Boca Raton, FL November, 2018; and abstract published in Innovations in Clinical Neuroscience, November-December 2018;15(11-12,suppl):S10.https://content.equisolve.net/tonixpharma/media/1d0c4055b2863fc74e1ef45f9ddaf42b.pdf

CAPS-5=Clinician-Administered PTSD Scale for DSM-5; MCFB=mean change from baseline; mITT=modified Intent-to-Treat sample; TST=time since trauma
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Summary of Clinical Experience with Tonmya/ TNX-102 SL in PTSD

Median time since trauma (TST) in TNX-102 SL 5.6 mg group in the P301/HONOR study (9.5 years) was longer than P201/AtEase study (6 years)

28

- · Both studied military-related PTSD
- · Time has passed since the surge in Iraq

In retrospective analysis, the \leq 9 year subgroup of P301 study had similar results as the P201 study (primary and secondary)

- · TST is important in placebo-controlled clinical study
- Potential enrichment in ≤ 9 years TST subgroup for treatment responders

The ≤ 9 year subgroup of P301 may be enriched for "Remitting Phase" of PTSD¹⁻⁴

· Expect remitting phase of PTSD is more amenable to drug studies

Results from retrospective analyses lead to improved Phase 3 study design

Kessler et al. Arch Gen Psychiatry 1995;52:1048-1060. Varmenta et al. BMC Psychiatry 2018;18:48. TGalatzer-Lavy et al. PLOS OWE 2013;8:e70084. Perkonigg et al. Am J Psychiatry 2005;162:1320-1327.



New Phase 3 P302/RECOVERY Study – Expected to Start 1Q 2019



General study characteristics: Primary endpoint: Randomized, double-blind, placebo-controlled study with CAPS-5¹ mean change from baseline at Week 4 (Tonmya 5.6 mg vs. placebo) . baseline CAPS-5¹ ≥ 33 in approximately 30 U.S. sites · Enrollment restricted to study participants with PTSD who experienced an index trauma \leq 9 years from the date of Key Secondary endpoints include: screening · CAPS-5 mean change from baseline at Week 12 (Tonmya 5.6 mg · Both civilian and military-related PTSD to be included vs. placebo) · Change from baseline Clinical Global Impression - Severity scale Tonmya once-daily at bedtime · Change from baseline Sheehan Disability Scale total score Potential pivotal efficacy study to support NDA approval Placebo once-daily at bedtime 12 weeks -≁ Primary endpoint CAPS-51 at Week 4 CAPS-5 = Clinician-Administered PTSD Scale for DSM-5 @ 2019 Tonix Pharmaceuticals Holding Corp.



Late-Stage PTSD Drug Candidates

Tonmya

 Phase 3 development focused on military-related and civilian PTSD; showed activity in treatment of military-related PTSD in large multi-center trials 30

MDMA-assisted psychotherapy

 Breakthrough therapy that is Phase 3-ready; showed activity in a Phase 2 study of PTSD; enrolling in Phase 3

Other drugs currently (or recently) in Phase 2 development

- Rexulti® (brexpiprazole) Otsuka/Lundbeck; atypical antipsychotic; positive clinical results from Phase 2 study reported in November 2018 for brexpiprazole, when used in combination with an approved PTSD medication, sertraline, but not as monotherapy
- NYX-783 Aptinyx; NMDA receptor modulator (enrolling for 8-week Phase 2 study of 144 patients using 50 mg either once daily or once weekly)
- BNC-201 Bionomics; nicotinic receptor modulator (program planned to resume after reformulation)



TNX-102 SL – Bedtime Treatment for Multiple Potential Indications



Management of Fibromyalgia (FM) – chronic pain condition

- TNX-102 SL studied at low dose (2.8 mg) half the dose being developed for PTSD – did not separate from placebo on primary endpoint, average pain improvement (responder analysis)
- Retrospective analysis showed average pain improvement (secondary endpoint) after 12 weeks of treatment showed statistical significance (P< 0.05, MMRM)
- Low dose TNX-102 SL (2.8 mg) showed an improvement in sleep quality in Phase 2 and Phase 3 FM trials
- Efficacy of TNX-102 SL 5.6 mg in FM can be studied in a potential pivotal study to support product registration

Agitation in Alzheimer's Disease

- Fast Track designation granted July 2018
- Phase 2/ potential pivotal efficacy study protocol received FDA comments in October 2018





FDA designated Fast Track development program

Significant unmet need

• No FDA approved drugs for the treatment of agitation in Alzheimer's

Mechanism of improving sleep quality

· Sleep disturbance is a significant and common symptom in Alzheimer's

Pharmacological advantages outweigh potential concerns of using TNX-102 SL in treating agitation in Alzheimer's disease

Blocks 3 receptors, not just one (e.g., 5-HT_{2A})

TNX-102 SL for Agitation in Alzheimer's – Regulatory Status and Registration Strategy



FDA confirmed no additional study was needed prior to IND submission

 Pre-IND meeting established open dialogue with the FDA on pivotal clinical study design and efficacy endpoints to support product registration

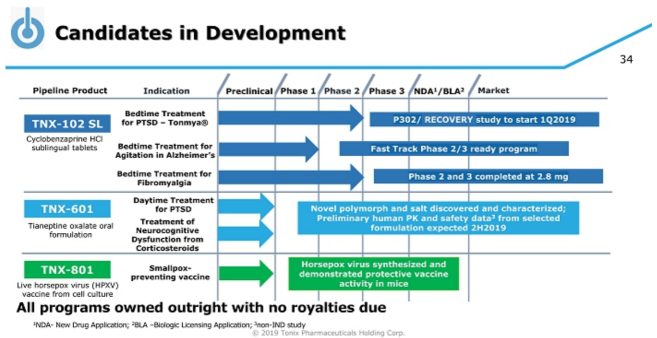
Proposed Phase 2 IND study can potentially serve as a pivotal efficacy study to support NDA approval

· FDA comments on final protocol received October 2018

Registration Strategy of TNX-102 SL for agitation in Alzheimer's disease

 Efficacy Supplement (sNDA¹) may be leveraged from the PTSD/FM development program and supported by Initial NDA approval for PTSD/FM

Supplemental New Drug Application



OVERTIMATE OT A Set UP: A Potential Clinical Candidate for PTSD

Pre-IND Candidate	 Targeted as a 1st line monotherapy for PTSD: oral formulation for daytime dosing Leverages expertise in PTSD (clinical and regulatory experience, market analysis, etc.) Mechanism of Action (MOA) is different from TNX-102 SL Tianeptine sodium (amorphous) has been approved in EU, Russia, Asia and Latin America for depression since 1987 with established post-marketing experience Identified new oxalate salt polymorph with improved pharmaceutical properties ideal for reformulation Preliminary human pharmacokinetic and safety data (non-IND study) from selected formulation expected in second half 2019 					
Targeting a	 Issued patent on steroid-induced cognitive impairment and memory loss issues 					
Condition with Significant	 Clinical evidence for PTSD Several studies have shown tianeptine to be active in the treatment of PTSD¹⁻⁴ 					
Unmet Need						
 Frančákovič T, et al. Psychiatr Danub. 2011 S Romyantseva GM and, Stepanov AL. Neurosci Ateksandrovski TA, et al. Zh Neurol Psikhiatr I Onder E, et al. Eur Psychiatry. 2006 (3):174-9 	Behav (hysiol. 2008 Jan;38(1):55-61. pMID: 18097761 m 5 5 Korsakova. 2005;156(11):24-0. PMID: 16329631 [Russian]					

35



TNX-801 (Synthesized Live Horsepox Virus): A Smallpox-Preventing Vaccine Candidate

Pre-IND Stage	 Potential improvement over current biodefense tools against smallpox ✓ Leverages Tonix's government affairs effort ✓ Collaboration with Professor David Evans and Dr. Ryan Noyce at University of Alberta ✓ Demonstrated protective vaccine activity in mice ✓ Patent application on novel vaccine submitted Regulatory strategy
Pre-Ind Stage	 We intend to meet with FDA to discuss the most efficient and appropriate investigational plan to support the licensure, either: ✓ Application of the "Animal Rule", or ✓ Conducting an active comparator study using ACAM2000 Good Manufacturing Practice (GMP) viral production process in development
Targeting a Potential Public Health Issue	 Material threat medical countermeasure under 21st Century Cures Act Qualifies for Priority Review Voucher (PRV) upon licensure* ✓ PRVs have no expiration date, are transferrable and have sold for ~\$125 M

36

*BLA/NDA priority 6-month review is expected.



NASDAQ: TNXP

Cash and cash equivalents, September 30, 2018	\$14.8 million
Net proceeds from equity offerings in 4Q2018	\$17.3 million
Common Stock outstanding post December 2018 underwritten equity offering (as of March 1, 2019)	4.7 million
Pro Forma Common Stock outstanding post December 2018 underwritten equity offering ¹ (as of March 1, 2019)	6.1 million

37

¹ Pro forma to include the remaining 1.4 million shares of Common Stock, not yet converted as of March 1, 2019, issuable upon conversion of the Series A Convertible Preferred Stock, as per terms of the December 2018 underwritten offering.



Management Team





Board of Directors



Seth Lederman, MD	Donald Landry, MD, PhD		
Chairman	Chair of Medicine, Columbia University		
Margaret Smith Bell Standard Life Investments, Putnam Investments, State Street Research	Adeoye "Oye" Olukotun, MD Squibb, BMS, Mallinckrodt, Esperion		
Patrick Grace	John Rhodes		
(qp) global family offices, Grace Institute	Chair, NYS Public Service Commission, CEO		
Foundation, WR Grace, Chemed	NYS Dept. of Public Service, Booz Allen		
Gen. David Grange (US Army, ret.)	James Treco		
Pharm-Olam, PPD, McCormick Foundation	First Chicago, Salomon Brothers/Citigroup		

b Milestones – Recently Completed and Upcoming

ď	July 2018	Completed P301/HONOR study interim analysis - result did not support study continuation but strengthened new Phase 3 study
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প্র	November 2018	Received FDA minutes confirming agreement on the design of P302/RECOVERY study
	First Quarter 2019	P302/RECOVERY study to be initiated
	First Quarter 2019	FM FDA Clinical Guidance meeting
	Second Half 2019	Preliminary human pharmacokinetic and safety data (non-IND study) from selected TNX-601 (tianeptine oxalate) formulation expected
	First Half 2020	Topline data from P302/RECOVERY study expected

40



Summary



Phase 3 development of new bedtime treatment for PTSD, including militaryrelated PTSD

- · Major unmet need; ~12 million Americans annually
- · Benefited from FDA 505(b)(2) NDA approval requirement

Complimentary day-time PTSD treatment in development

 Leverages development expertise in PTSD, i.e., regulatory, trial recruitment and execution

New indication in development for agitation in Alzheimer's Disease

- · Unmet medical need, no approved drug available
- Fast Track Phase 2/3 ready program

Fibromyalgia bedtime treatment in development

· IND ready to support Phase 3 potential pivotal efficacy study

Innovative vaccine in development to prevent Smallpox

- · Opportunity to supply stockpiling requirement; short development path
- · Studies in mice suggest improved safety profile







Exhibit 99.03





March 2019

Version P0159 3-5-19 (Doc 0437)



Cautionary Note on Forward-Looking Statements

2

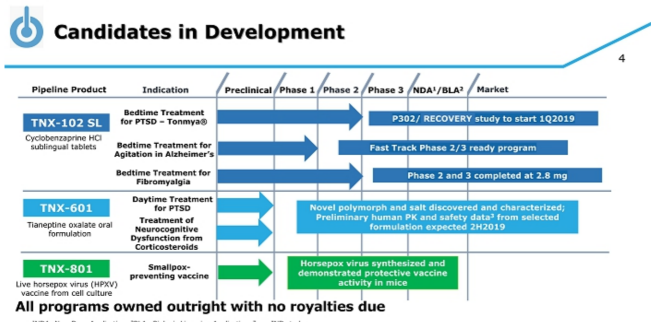
Certain statements in this presentation regarding strategic plans, expectations and objectives for future operations or results are "forward-looking statements" as defined by the Private Securities Litigation Reform Act of 1995. These statements may be identified by the use of forward-looking words such 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, substantial competition; 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 risks related to failure to obtain U.S. Food and Drug Administration clearances or approvals and noncompliance with its regulations. 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, 2017, as filed with the Securities and Exchange Commission (the "SEC") on March 9, 2018, and periodic reports and current reports filed with the SEC on or after the date thereof. All of Tonix's forwardlooking statements are expressly qualified by all such risk factors and other cautionary statements.



Posttraumatic Stress Disorder (PTSD) - Lead program; new bedtime Cyclobenzaprine Sublingual Tablets treatment - Tonmya®1 **INX-102 SI** P302/RECOVERY Phase 3 clinical study with Week 4 primary endpoint to initiate in 1Q2019 · Results from 2 efficacy studies improve the new Phase 3 study design New Phase P302/RECOVERY study design features accepted by the FDA² Agitation in Alzheimer's disease (AAD) IND³ ready to support Phase 2 potential pivotal efficacy study Fibromyalgia Syndrome (FM) IND³ ready to support Phase 3 potential pivotal efficacy study TNX-601⁴ – Daytime PTSD treatment and treatment of neurocognitive Pipeline dysfunction from corticosteroids Pre-IND candidate; nonclinical development ongoing TNX-801⁵ - Smallpox-preventing vaccine candidate Efficacy demonstrated in mouse model ¹ Tonmya has been conditionally accepted by the U.S. FDA as the proposed trade name for TNX-102 SL (cyclobenzaprine HCI sublingual tablets) for the treatment of PTSD. TNX-102 SL is an investigational new drug and has not been approved for any indication. ² FDA Meeting Minutes (November 26, 2018) ³ IND- Investigational New Drug Application ⁴ Timeetine evaluate

3

⁴ Tianeptine oxalate ⁵ Synthesized live horsepox virus



¹NDA- New Drug Application; ²BLA -Biologic Licensing Application; ³non-IND study (© 2019 Tonix Pharmaceuticals Holding Corp.



First investigational new drug to show treatment effect in military-related PTSD in two potential pivotal efficacy studies

 Phase 2 study (P201/AtEase) showed Tonmya 5.6 mg had a strong signal of treatment effect at Week 12 as measured by CAPS-5¹

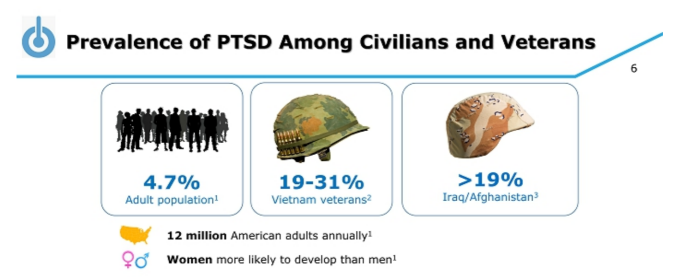
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- Phase 3 study (P301/HONOR) provided evidence of effectiveness as early as 4 weeks after treatment but diminished over time due to high placebo response
 - Retrospective analysis showed persistent effectiveness at Week 12 in subgroup with trauma ≤9 years from screening
- Both studies can be used as supportive evidence of efficacy and safety for Tonmya NDA submission
- No serious or unexpected adverse events related to Tonmya were reported

FDA feedback and acceptance on new Phase 3 study (P302/RECOVERY) received in November²

¹ CAPS-5 = Clinician-Administered PTSD Scale for DSM-5 ² FDA Meeting Minutes, November 26, 2018

٠



³Goldstein et al., 2016 (adjusted for 2019); ²Norris, PTSD Res Quar. 2013; ³Analysis of VA Health Care Utilization among Operation Enduring Freedom, Operation Iragi Freedom, and Operation New Dawn Veterans, from 1st Qtr FY 2002 through 2nd Qtr FY 2015, Washington, DC; Among 1.9M separated OEF/OIF/OND veterans, 1.2M have obtained VA healthcare; 685k evaluated by VA with possible mental disorder, and 379k diagnosed with PTSD.



Unmet Need for Effective and Safe Therapies for Treatment of Military PTSD

7

PTSD is signature wound of last 25 years of war

- · Affects servicemember health and performance, force readiness, retention
- · Believed to be the underlying cause of suicide in many cases

No FDA-approved products for PTSD since Pfizer's Zoloft[®] (sertraline) and GSK's Paxil[®] (paroxetine) circa 2000

- Neither has shown efficacy in military-related PTSD
- Male PTSD patients often unresponsive or intolerant of current treatments
- Side effects relating to sexual dysfunction, sleep and weight gain are commonly reported

U.S. Department of Defense (DoD) is working to understand and treat PTSD

- · Increased scrutiny of PTSD-related discharges for behavioral problems
- Wider recognition that PTSD is a service-related disability
- Collaboration with Army: Tonix-USAMMDA CRADA signed in 2015



TNX-102 SL Intellectual Property – U.S. Protection until 2034



- United States Patent and Trademark Office (USPTO) issued U.S. Patent No. 9,636,408 in May 2017 U.S. Patent No. 9,956,188 in May 2018 and U.S. Patent No. 10,117,936 in November 2018
- Japanese Patent Office (JPO) issued Japanese Patent No. 6310542 in March 2018
- New Zealand Intellectual Property Office (NZIPO) issued New Zealand Patent No. 631152 in May 2017

8

37 patent applications pending (2 allowed (US and South Africa))

Pharmacokinetics (PK) : Protection expected to 2033

- JPO issued Japanese Patent No. 6259452 in December 2017
- NZIPO issued New Zealand Patent No. 631144 in March 2017
- Taiwanese Intellectual Property Office issued Taiwanese Patent No. I590820 in July 2017
- 21 patent applications pending (1 allowed (Australia))

Method of use for active ingredient cyclobenzaprine : Protection expected to 2030

- USPTO issued U.S. Patent 9,918,948 in March 2018
- European Patent Office issued European Patent No. 2 501 234B1 in September 2017 (validated in 38 countries). Opposition filed in June 2018
- · 2 patent applications pending



Potential Therapeutic Advantages of Tonmya

Tonmya is believed to treat PTSD by improving sleep quality

- · The brain naturally processes memories during sleep
- PTSD sufferers' emotionally charged memories disturb sleep and disrupt the natural processing of memories during sleep
- Tonmya is believed to normalize memory processing and facilitate extinction consolidation (breaking the link between "triggers" and PTSD symptoms)

Tonmya is NEITHER a benzodiazepine nor a narcotic

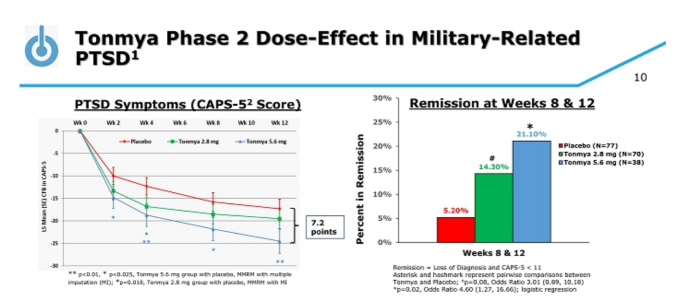
 The active ingredient of Tonmya, cyclobenzaprine, does <u>NOT</u> interact with the same receptors as traditional hypnotic sleep drugs associated with retrograde amnesia; is <u>NOT</u> an opiate

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Tonmya is non-addictive

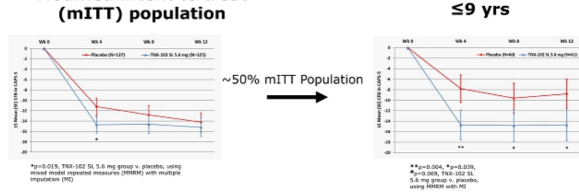
- Cyclobenzaprine is the active ingredient of an orally ingested immediate release tablet (Flexeril[®]), approved 40 years ago
- Flexeril's current labeling indicates no abuse and dependence concern at higher doses than Tonmya (15-30 mg/day v. 5.6 mg/day); NDA can be filed without drug abuse and dependency assessment studies

Once-daily sublingual dose taken at bedtime enhances patient adherence



³Completed Phase 2 P201/AtEase study: Retrospective analysis of Tonmya 5.6 mg on CAPS-5 ≥33 (high-moderate) subgroup. Primary analysis of P201/AtEase was on Tonmya 2.8 mg in participants with entry CAPS-5 ≥29, moderate PTSD severity. ³Clinician administered PTSD Scale for DSM-5

Primary Outcome (CAPS-5) in Phase 3 (mITT) and ≤9 Years Time Since Trauma (TST) Subgroups Phase 3 P301/HONOR Study¹ Modified intent to treat

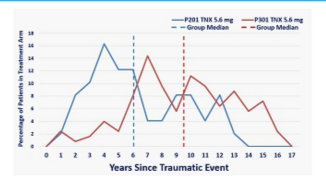


¹Phase 3 P301/HONOR study: stopped in July 2018. Separation on primary endpoint did not cross pre-specified study continuation threshold at Week 12 in the interim analysis at ~50% randomization; no safety or tolerability issues discovered.



Retrospective Comparison of Time Since Trauma in P201/AtEase versus P301/HONOR (Tonmya 5.6 mg Groups)

12



P301 study was initiated approximately two years later than Phase 2 P201

 The median time since trauma in Phase 3 was 9.5 years compared to the median time since trauma in Phase 2 of 6.0 years for TNX-102 SL 5.6 mg treated groups

6

Sustained Remission in Phase 2 and Phase 3 Studies: Retrospective Analyses of P201 Entry CAPS-5 ≥33 and P301 ≤9 Years Since Trauma Subgroups

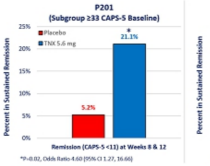
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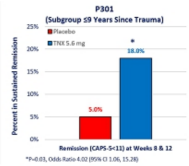
Remission is a clinical state that is essentially asymptomatic

In order to confirm remission:

 Determined rates of participants who met remission status at both Week 8 and Week 12

Rate of remission in ≤9 years since trauma group in P301 is similar to baseline CAPS-5 ≥ 33 group in P201¹





¹Majority of P201 participants were ≤ 9 years since trauma and ~80% of P201 participants and all of P301 participants were ≥ 33 CAPS-5 at baseline



Adverse Events (AEs) in P201/AtEase and P301/HONOR Studies

	P201		P301		
Category of Adverse Reaction Preferred Term	Placebo (N=94)	TNX 2.8 mg (N=93)	TNX 5.6 mg (N=50)	Placebo (N=134)	TNX 5.6 mg (N=134)
Systemic Adverse Events**					
Somnolence	6.4%	11.8%	16.0%	9.0%	15.7%
Dry mouth	10.6%	4.3%	16.0%		
Headache	4.3%	5.4%	12.0%		
Insomnia	8.5%	7.5%	6.0%		
Sedation	1.1%	2.2%	12.0%		
Local Administration Site Reactions* [#]					
Hypoaesthesia oral	2.1%	38.7%	36.0%	1.5%	37.3%
Paraesthesia oral	3.2%	16.1%	4.0%	0.7%	9.7%
Glossodynia	1.1%	3.2%	6.0%		
Product Taste Abnormal				3.0%	11.9%

14

²only adverse events (AEs) are listed that are at a rate of \geq 5% in any TNX-treated group *no values in a row for either study means the AE in the active group(s) in that study was at a rate of <5%

No serious or unexpected AEs in P201 or P301 related to Tonmya

- Systemic AEs comparable between studies and also consistent with those described in approved oral cyclobenzaprine product labeling
- Severity and incidence of oral hypoesthesia (oral numbness) are not dose related and similar in both studies
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Response to Tonmya for Female Participants in P301/HONOR Study¹



Females made up only 11% of the P301/HONOR study mITT population

Difference in mean change from baseline in CAPS-5 in females between placebo (N=17) and Tonmya 5.6 mg (N=10) was:

- At 4 weeks -11.5 points
- · At 12 weeks -9.1 points

Indicates substantial separation from placebo in the small number of female participants

Predicts therapeutic response to Tonmya 5.6 mg likely in mixed civilian and military PTSD population to be studied in upcoming P302/RECOVERY trial

Civilian PTSD population tends to be about 2/3 female

¹Time Since Trauma in PTSD: Phase 3 Multi-Center, Double-Blind, Placebo-Controlled Trial of TNX-102 SL, a Sublingual Formulation of Cyclobenzaprine, in Military-Related PTSD (Study TNX-CY-P301) Presented at CNS Summit in Boca Raton, FL November 1-4, 2018 and abstract published in Innovations in Clinical Neuroscience, November-December 2018;15(11-12,suppl):S10.<u>https://content.eguisolve.net/tonixpharma/media/1d0c4055b2863fc74e1ef45f9ddaf42b.pdf</u>

Response to Tonmya for Non-Combat Traumas in P301/HONOR Study in ≤9 Years Time Since Trauma Subgroup¹

Non-combat traumas studied are similar to traumas experienced in civilian populations with PTSD

To determine the therapeutic effects of Tonmya 5.6 mg in a mixed civilian and military population, difference in MCFB in CAPS-5 was assessed in non-combat traumas in \leq 9 years TST subgroup (placebo N=14, Tonmya 5.6 mg N=10):

16

- At 4 weeks -4.8 points
- At 12 weeks -4.4 points

Non-combat traumas treated with Tonmya 5.6 mg showed clinically meaningful separation from placebo at Weeks 4 and 12, suggesting a mixed civilian and military sample within 9 years of index trauma may show a therapeutic response to Tonmya

¹Time Since Trauma in PTSD: Phase 3 Multi-Center, Double-Blind, Placebo-Controlled Trial of TNX-102 SL, a Sublingual Formulation of Cyclobenzaprine, in Military-Related PTSD (Study TNX-CY-P301) Presented at CNS Summit in Boca Raton, FL November, 2018 and abstract published in Innovations in Clinical Neuroscience, November-December 2018;15(11-12,suppl):S10.https://content.equisolve.net/tonixpharma/media/1d0c4055b2863fc74e1ef45f9ddaf42b.pdf

CAPS-5=Clinician-Administered PTSD Scale for DSM-5; MCFB=mean change from baseline; mITT=modified Intent-to-Treat sample; TST=time since trauma
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New Phase 3 P302/RECOVERY Study – Expected to Start 1Q 2019



General study characteristics: Primary endpoint: Randomized, double-blind, placebo-controlled study with CAPS-5¹ mean change from baseline at Week 4 (Tonmya 5.6 mg vs. placebo) . baseline CAPS-5¹ ≥ 33 in approximately 30 U.S. sites · Enrollment restricted to study participants with PTSD who experienced an index trauma \leq 9 years from the date of Key Secondary endpoints include: screening · CAPS-5 mean change from baseline at Week 12 (Tonmya 5.6 mg · Both civilian and military-related PTSD to be included vs. placebo) · Change from baseline Clinical Global Impression - Severity scale Tonmya once-daily at bedtime · Change from baseline Sheehan Disability Scale total score Potential pivotal efficacy study to support NDA approval Placebo once-daily at bedtime 12 weeks -≁ Primary endpoint CAPS-51 at Week 4 CAPS-5 = Clinician-Administered PTSD Scale for DSM-5 @ 2019 Tonix Pharmaceuticals Holding Corp.



Late-Stage PTSD Drug Candidates

Tonmya

 Phase 3 development focused on military-related and civilian PTSD; showed activity in treatment of military-related PTSD in large multi-center trials 18

MDMA-assisted psychotherapy

 Breakthrough therapy that is Phase 3-ready; showed activity in a Phase 2 study of PTSD; enrolling in Phase 3

Other drugs currently (or recently) in Phase 2 development

- Rexulti® (brexpiprazole) Otsuka/Lundbeck; atypical antipsychotic; positive clinical results from Phase 2 study reported in November 2018 for brexpiprazole, when used in combination with an approved PTSD medication, sertraline, but not as monotherapy
- NYX-783 Aptinyx; NMDA receptor modulator (enrolling for 8-week Phase 2 study of 144 patients using 50 mg either once daily or once weekly)
- BNC-201 Bionomics; nicotinic receptor modulator (program planned to resume after reformulation)



TNX-102 SL – Bedtime Treatment for Multiple Potential Indications



Management of Fibromyalgia (FM) – chronic pain condition

- TNX-102 SL studied at low dose (2.8 mg) half the dose being developed for PTSD – did not separate from placebo on primary endpoint, average pain improvement (responder analysis)
- Retrospective analysis showed average pain improvement (secondary endpoint) after 12 weeks of treatment showed statistical significance (P< 0.05, MMRM)
- Low dose TNX-102 SL (2.8 mg) showed an improvement in sleep quality in Phase 2 and Phase 3 FM trials
- Efficacy of TNX-102 SL 5.6 mg in FM can be studied in a potential pivotal study to support product registration

Agitation in Alzheimer's Disease

- Fast Track designation granted July 2018
- Phase 2/ potential pivotal efficacy study protocol received FDA comments in October 2018

TNX-102 SL for Agitation in Alzheimer's – Regulatory Status and Registration Strategy



FDA confirmed no additional study was needed prior to IND submission

 Pre-IND meeting established open dialogue with the FDA on pivotal clinical study design and efficacy endpoints to support product registration

Proposed Phase 2 IND study can potentially serve as a pivotal efficacy study to support NDA approval

FDA comments on final protocol received October 2018

Registration Strategy of TNX-102 SL for agitation in Alzheimer's disease

 Efficacy Supplement (sNDA¹) may be leveraged from the PTSD/FM development program and supported by Initial NDA approval for PTSD/FM

Supplemental New Drug Application

OVERTIMATE OT A POTENTIAL OT A POTENTIAL CLANCED A POTENTIAL CLINICAL CANDIDATE FOR PTSD

	Targeted as a 1 st line monotherapy for PTSD: oral formulation for daytime dosing	
	 Leverages expertise in PTSD (clinical and regulatory experience, market analysis, etc.) 	
	Mechanism of Action (MOA) is different from TNX-102 SL	
Pre-IND Candidate	 Tianeptine sodium (amorphous) has been approved in EU, Russia, Asia and Latin America for depression since 1987 with established post-marketing experience 	
	 Identified new oxalate salt polymorph with improved pharmaceutical properties ideal for reformulation 	
	 Preliminary human pharmacokinetic and safety data (non-IND study) from selected formulation expected in second half 2019 	
	Filed patent application on novel salt polymorph	
Targeting a	 Issued patent on steroid-induced cognitive impairment and memory loss issues 	
Condition with Significant	Clinical evidence for PTSD	
Unmet Need	 Several studies have shown tianeptine to be active in the treatment of PTSD¹⁻⁴ 	
 Frančšković T, et al. Psychiatr Danub. 2011 Sep;23(3):257-63. MMID: 21963693 Pournyantseva GM and, Stopanov AL, Neurosci Behav Mysiol. 2008 San;38(1):55-61. PMID: 18097761 Aleksandrovski TJ, et al. Zh. Novrol Bishiaru Im S 5 Korsakova. 2005;105(11):24-9. PMID: 16329631 [Russian] Onder E, et al. Eur Psychiatry. 2006 (3):174-9. PMID: 15964747 		

21



TNX-801 (Synthesized Live Horsepox Virus): A Smallpox-Preventing Vaccine Candidate

	Potential improvement over current biodefense tools against smallpox ✓ Leverages Tonix's government affairs effort ✓ Collaboration with Professor David Evans and Dr. Ryan Noyce at University of Alberta ✓ Demonstrated protective vaccine activity in mice
Pre-IND Stage	 Patent application on novel vaccine submitted Regulatory strategy
	 We intend to meet with FDA to discuss the most efficient and appropriate investigational plan to support the licensure, either:
	✓ Application of the "Animal Rule", or ✓ Conducting an active comparator study using ACAM2000
	Good Manufacturing Practice (GMP) viral production process in development
Targeting a	Material threat medical countermeasure under 21st Century Cures Act
Potential Public Health Issue	 Qualifies for Priority Review Voucher (PRV) upon licensure* ✓ PRVs have no expiration date, are transferrable and have sold for ~\$125 M

22

*BLA/NDA priority 6-month review is expected.



NASDAQ: TNXP

Cash and cash equivalents, September 30, 2018	\$14.8 million
Net proceeds from equity offerings in 4Q2018	\$17.3 million
Common Stock outstanding post December 2018 underwritten equity offering (as of March 1, 2019)	4.7 million
Pro Forma Common Stock outstanding post December 2018 underwritten equity offering ¹ (as of March 1, 2019)	6.1 million

23

¹ Pro forma to include the remaining 1.4 million shares of Common Stock, not yet converted as of March 1, 2019, issuable upon conversion of the Series A Convertible Preferred Stock, as per terms of the December 2018 underwritten offering.



Management Team



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