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): August 7, 2023

TONIX PHARMACEUTICALS HOLDING CORP.

(Exact name of registrant as specified in its charter)

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

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

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

Check the appropriate box below if the Form 8-K filing is intended to simultaneously satisfy the filing obligation of the registrant under any of the following provisions (see

General Instruction A.2. below	r):	
☐ Soliciting material pursuan ☐ Pre-commencement comm	to Rule 14a-12 under the Securities Act (17 CFR 240 to Rule 14a-12 under the Exchange Act (17 CFR 240 unications pursuant to Rule 14d-2(b) under the Exchanunications pursuant to Rule 13e-4(c) under the Exchanunications (a) and (b) of the Act:	.14a-12) ge Act (17 CFR 240.14d-2(b))
Title of each class	Trading Symbol(s)	Name of each exchange on which registered
Common Stock	TNXP	The NASDAQ Capital Market
	ny, indicate by check mark if the registrant has elected pursuant to Section 13(a) of the Exchange Act. □	not to use the extended transition period for complying with any new or revised financial
Tonix Pharmaceutica and at investor conferences, as	nd which the Company intends to place on its website	r presentation, which is used to conduct meetings with investors, stockholders and analysts, which may contain nonpublic information. A copy of the presentation is filed as Exhibit
99.01 hereto and incorporated The information in th	·	uding Exhibit 99.01 attached hereto, shall not be deemed "filed" for purposes of Section 18

Item 9.01 Financial Statements and Exhibits.

(d)	Exhibit	
	No.	Description.
·	<u>99.01</u>	Corporate Presentation by the Company for August 2023
	104	Cover Page Interactive Data File (embedded within the Inline XBRL document)

of the United States Securities Exchange Act of 1934 (the "Exchange Act") or otherwise subject to the liabilities of that section, nor shall they be deemed incorporated by reference in any filing under the United States Securities Act of 1933 or the Exchange Act, except as shall be expressly set forth by specific reference in such a filing.

SIGNATURE

TONIX PHARMACEUTICALS HOLDING CORP.

Date: August 7, 2023 By: /s/ Bradley Saer

By: /s/ Bradley Saenger Bradley Saenger Chief Financial Officer



Cautionary Note on Forward-Looking Statements

Certain statements in this presentation regarding strategic plans, expectations and objectives for future operations or results are "forward-looking statements" as defined by the Private Securities Litigation Reform Act of 1995. These statements may be identified by the use of forward-looking words such as "anticipate," "believe," "forecast," "estimate" and "intend," among others. These forward-looking statements are based on Tonix's current expectations and actual results could differ materially. There are a number of factors that could cause actual events to differ materially from those indicated by such forward-looking statements. These factors include, but are not limited to, the risks related to failure to obtain FDA clearances or approvals and noncompliance with FDA regulations; risks related to the failure to successfully market any of our products; risks related to the timing and progress of clinical development of our product candidates; our need for additional financing; uncertainties of patent protection and litigation; uncertainties of government or third party payor reimbursement; limited research and development efforts and dependence upon third parties; and substantial competition. As with any pharmaceutical under development, there are significant risks in the development, regulatory approval and commercialization of new products. The forward-looking statements in this presentation are made as of the date of this presentation, even if subsequently made available by Tonix on its website or otherwise. Tonix does not undertake an obligation to update or revise any forward-looking statement, except as required by law. Investors should read the risk factors set forth in the Annual Report on Form 10-K for the year ended December 31, 2022, as filed with the Securities and Exchange Commission (the "SEC") on March 13, 2023, and periodic reports and current reports filed with the SEC on or after the date thereof. All of Tonix's forward-looking statements are expressly qualified by all such risk factors and other cautionary

TÖNIX 2

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Investment Highlights



DIVERSE PIPELINE

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



IN-HOUSE CAPABILITIES

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



STRATEGIC PARTNERSHIPS

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



FINANCIAL POSITION

Tonix announced pricing of a \$7 M financing on July 27, 2023. Tonix has no debt.

TONIX PHARMACEUTICALS

Pipeline: Key Clinical Development Programs

Candidates*	Indication	Status/Next Milestone	
TNX-102 SL ¹	Fibromyalgia (FM) Long COVID (PASC²)	Mid-Phase 3 – enrollment complete Phase 2 – enrollment complete	
TNX-1300 ³	Cocaine Intoxication - FDA Breakthrough Designation	Mid-Phase 2, Targeted 3Q 2023 Start	
TNX-1900 ⁴	Prevention of Chronic Migraine	Phase 2 – enrollment complete ⁵	
TNX-601 ER	Depression Pha	Phase 2 – enrollment complete ⁶	
TNX-2900 ⁷	2900 ⁷ Prader-Willi Syndrome - FDA Orphan Drug Designation Phase 2 re		
TNX-1500 ⁸	Organ Transplant Rejection/ Autoimmune Conditions	Phase 1, Targeted 3Q 2023 Start	

'All of Tonix's product candidates are investigational new drugs or biologics and none has been approved for any indication.

'TNX-102 SL (cyclobenzaprine HCI sublingual tablets) also has active INDs for Agitation in Alzheimer's Disease (AAD), Alcohol Use Disorder (AUD), and Posttraumatic Stress Disorder (PTSD). All indications are Phase 2 ready.

'Post-Acute Sequelae of COVID-19.

'PIXX-1300 (double-mutant coaline esterase) is licensed from Columbia University.

'Acquired from Trigernins, license agreement with Stanford University, Planned investigator-initiated Binge Eating Disorder (BED) study is expected start 2Q 2023.

'A Phase 2 trial under an investigator-initiated IND has been completed in the U.S. using TNX-1900

'Phase 1 trial for formulation development was completed outside of the U.S.: Other potential indications include PTSD and neurocognitive dysfunction from steroids

'Co-exclusive locense agreement with French National Institute of Health and Medical Research (Inserm)

*anti-CD40L humanized monoclonal antibody – IND cleared

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Late-Stage CNS Programs¹

Four Studies Expecting Topline in the Next Three Quarters (by End of 1Q24)

Active Studies

- 23Q3 Topline:
 - Proof-of-Concept TNX-102 SL for fibromyalgia-type Long COVID (enrollment complete)
 - Clinical Phase of trial complete
- 23Q4 Topline:
 - TNX-1900 for migraine headache (enrollment complete) P2 Proof-of-Concept TNX-601 ER for depression (enrollment complete) P2 Proof-of-Concept
 - Potential NDA enabling P3 TNX-102 SL for fibromyalgia (enrollment complete)

Entering Phase 2

- In 3Q 2023:
 - TNX-1300 for cocaine intoxication (FDA Breakthrough Therapy)

Potential Pivotal Study

P2

¹Not approved for any indication



Two Marketed Proprietary Migraine Drugs Autoinjector and Nasal Spray (Non-oral) Formulations of Sumatriptan

Zembrace® SymTouch® (sumatriptan injection) 3 mg¹

Tosymra® (sumatriptan nasal spray) 10 mg²

- Each indicated for the treatment of acute migraine with or without aura in adults
- Sumatriptan remains the acute migraine 'gold standard' treatment for many patients and continues to represent the largest segment of the market in terms of unit sales3
- Each may provide migraine pain relief in as few as 10 minutes for some patients^{1,2,4,5}
- Each bypasses GI tract provides convenient administration
- Patents to 2036 (Zembrace) and 2031 (Tosymra)

Upsher-Smith Laboratories Providing Certain Commercial Operations

- Product acquisition closed on June 30, 2023
- To support the transition of the products, Upsher-Smith is providing certain commercial operations, regulatory and other transition services to Tonix Medicines for up to nine months after closing, in exchange for agreed upon service fees.

**Pembrace SymTouch [package insert], Maple Grove, MN: Upsher-Smith Laboratories, LLC: February 2021 - For more information, talk to your provider and read the Patient Information and Instructions for Use, — Important Safety Information is provided in the appendix.

**Tosymra [package insert], Maple Grove, MN: Upsher-Smith Laboratories, LLC: Feb 2021. For more information, talk to your provider and read the Patient Information and Instructions for Use, — Important Safety Information is provided in the appendix

**Upsher-Smith Laboratories, LLC; Data On File, 2023

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

**Wendt J, et al. A randomized, double-blind, placebo-controlled trial of the efficacy and tolerability of a 4-mg dose of subcutaneous sumatriptan for the treatment of acute migraine attacks in adults. Clinical Therapeutics. 2006;29(45):517-526.

**Tosymrate Safety Sa

Tonix Medicines is our Commercial Subsidiary

Marketed products for the treatment of acute migraine in adults with or without aura Zembrace® SymTouch® (sumatriptan injection) 3 mg1

Tosymra® (sumatriptan nasal spray) 10 mg²

Both are proprietary non-oral formulations of sumatriptan that bypass the gastrointestinal tract

Headed by President Jim Hunter

- Industry veteran experience in CNS products
- **Built Validus Pharmaceuticals**

Consolidated Product Sales for the 12 months ended March 31st 2023

- Factory sales: \$30.4M³
- Net sales: \$16.4M³

Retail Product Sales for the 12 months ended December 31st 2022

Retail sales: ~\$23 M (Zembrace ~\$19.6 M and Tosymra ~\$3.5 M)⁴

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

- Contract includes a transition period during which Tonix expects to secure its own contracts
- ¹Zembrace SymTouch [package insert]. Maple Grove, MN: Upsher-Smith Laboratories, LLC: February 2021 For more information, talk to your provider and read the Patient Information and Instructions for Use
- Important Safety Information is provided in the appendix

 Tosymra [package insert]. Maple Grove, MN: Upsher-Smith Laboratories, LLC: Feb 2021 . For more information, talk to your provider and read the Patient Information and Instructions for Use, Important Safety
- viated financial statements of assets acquired from Upsher-Smith Laboratories, LLC as filed in the 8-K/A dated July 18, 2023 https://fir.tonixpharma.com/sec-filings/all-sec-
- Paudited abbreviated financial statements of assets sugarities from the paner Samu. Specific Statements of assets sugarities from the SMART database estimates Zembrace sales of \$19.6 M and Tosymra sales of \$3.5 M © 2023 Tonix Pharmaceuticals Holding Corp.

Administration of Zembrace and Tosymra Bypass the GI Tract

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

Potential to provide a treatment option for migraines complicated by severe nausea and vomiting

Need for acute non-oral treatments

- GI absorption may be inconsistent in migraineurs due to gastric stasis (also called "gastroparesis")¹⁻⁴
- Nausea and vomiting are symptoms of migraine⁵

Existing intranasal products

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

New intranasal products bring attention to this non-oral route

- Pfizer's Zavzpret® (zavegepant), FDA approved in March, 2023¹ is the first intranasal gepant
- Impel NeuroPharma's Trudhesa® (dihydroergotamine) FDA approved 2021²
 - Precision Olfactory Delivery (POD) technology targets vascular-rich upper nasal space





Potential for Zembrace and Tosymra in Evolving Migraine Market

Documented efficacy of Zembrace^{1,2} and Tosymra³⁻⁵ as abortive treatments for acute migraine

- Migraine pain relief possible in as few as 10 minutes for some patients and convenient administration.
- Potential to address the unmet needs of patients using traditional or emerging oral acute migraine medications, particularly for rapid-onset treatment

Zembrace and Tosymra have potential as first-line or rescue medications

- Depending on patient need, prescribers may utilize to treat acute migraine either first-line or in the rescue position in a comprehensive migraine toolbox
- Fast onset of action, high pain-relief rates
- For certain patients who present to ERs, Zembrace and Tosymra also provide ER staff a straightforward treatment option

¹Zembrace SymTouch [package insert]. Maple Grove, MN: Upsher-Smith Laboratories, LLC: 2019.

²Landy, S. et al. Efficacy and safety of DFN-11 (sumatriptan injection, 3 mg) in adults with episodic migraine: a multicenter, randomized, double-blind, placebo-controlled study. *J Headache Pain*. 19, 69 (2018).

(2018).

Tosymra (package insert), Maple Grove, MN: Upsher-Smith Laboratories, LLC: Feb 2021.

Mathew NT, et al. Dose ranging efficacy and safety of subcutaneous sumatriptan in the acute treatment of migraine. US Sumatriptan Research Group. Arch Neurol. 1992.49(12):1271-1276.

Wendt J, et al. A randomized, double-blind, placebo-controlled trial of the efficacy and tolerability of a 4-mg dose of subcutaneous sumatriptan for the treatment of acute migraine attacks in a Clinical Therapeutics. 2006;29(4):517-526.

Intravail is a registered trademark of Aegis Therapeutics, LLC, a wholly owned subsidiary of Neurelis, Inc.





TNX-102 SL*



Cyclobenzaprine (Protectic®) Pipeline in a Product

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

Potent binding and antagonist activities at the serotonergic-5-HT2A, adrenergic-α1, histaminergic-H1, and muscarinic-M1 cholinergic receptors to facilitate restorative sleep

Innovative and proprietary PROTECTIC® Rapid drug exposure following nighttime administration

Differentiators:

Relative to Oral Cyclobenzaprine

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

Relative to Standard of Care

- Potential for better tolerability while maintaining efficacy
- Not scheduled with no recognized abuse potential

Patents Issued

INX-102 SL has not been approved for any indicatio

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Fibromyalgia

Status: Mid-Phase 3

- · One positive Phase 3 study (RELIEF) completed
- Second Phase 3 study (RALLY) missed primary endpoint
- · Confirmatory Phase 3 study (RESILIENT) enrollment complete

Next Steps: Topline results expected 4Q 2023

Fibromyalgia-Type Long COVID

Status: Phase 2

· Phase 2 study (PREVAIL) enrollment complete

Next Steps: Topline results expected 3Q 2023



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

PROFILE

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

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



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

DEVELOPMENT PROGRAM

Market Entry: Fibromyalgia

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

Status: One Positive Phase 3 study RELIEF

Second Phase 3 study RALLY missed primary endpoint

Confirmatory Phase 3 study RESILIENT enrollment complete

Next Steps: Topline results expected 4Q 2023

Patents Issued

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

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

Addedman et al. (2023) Arthritis Care & Research "Efficacy and Safety of TNX-102 SL (Sublingual Cyclobenzaprine) for the Treatment of Fibromyalgia: Results From the RELIEF Trial", doi: 10.1002/acr.25142. Epub ahead of print. PMID: 37165930.

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TNX-102 SL: Phase 3 RESILIENT Study Design

General study characteristics:

- · Randomized, double-blind, placebo-controlled study in fibromyalgia
- U.S. sites only, completed enrollment of 457 patients

Primary Endpoint:

- Daily diary pain severity score change from baseline to Week 14 (TNX-102 SL vs. placebo)
 - · Weekly averages of the daily numerical rating scale scores

Key Secondary Endpoints:

- Fibromyalgia Impact Questionnaire Revised (FIQ-R) Symptom Domain score
- Patient Global Impression of Change responder analysis
- FIQ-R Function Domain score
- PROMIS Sleep Disturbance instrument
- PROMIS Fatigue instrument
- Weekly average of the daily diary assessment of sleep quality

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

Placebo once-daily at bedtime

14 weeks

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

ClinicalTrials.gov Identifier: NCT05273749 A Phase 3 Study to Evaluate the Efficacy and Safety of TNX-102 SL Taken Daily in Patients With Fibromyalgia (RESILIENT)

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

PROFILE

- · Occurs in approximately 19% of recovered COVID-19 patients1
- · As many as 40% of Long COVID patients experience multi-site pain, a hallmark of fibromyalgia^{2,3}
- · Symptoms of Long COVID, like multi-site pain, fatigue and insomnia, are the hallmarks of chronic pain syndromes like fibromyalgia and myalgic encephalomyelitis/chronic fatigue syndrome (ME/CFS)
- · In August 2022, the HHS released the National Research Action Plan on Long COVID4 which endorses the connection between Long COVID and ME/CFS

DEVELOPMENT PROGRAM

Market Entry: Fibromyalgia-Type Long COVID (PASC)

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

Status: Phase 2 study PREVAIL enrollment complete

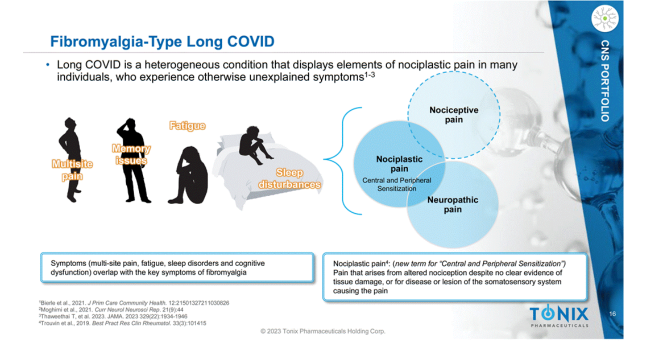
Next Steps: Topline results expected 3Q 2023

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

⁴Department of Health and Human Services, Office of the Assistant Secretary for Health. 2022. National Research Action Plan on Long COVID.







TNX-102 SL: Phase 2 PREVAIL Study Design

PREVAIL Study

Study characteristics:

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

Primary Endpoint:

- Daily diary pain severity score change from baseline to Week 14 (TNX-102 SL vs. placebo)
 - Weekly averages of the daily numerical rating scale scores

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

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

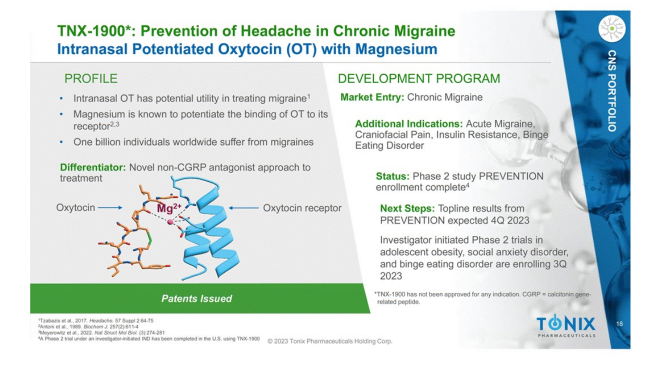
Placebo once-daily at bedtime

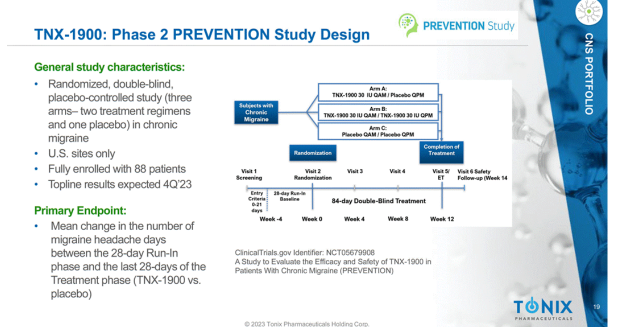
ClinicalTrials.gov Identifier: NCT05472090

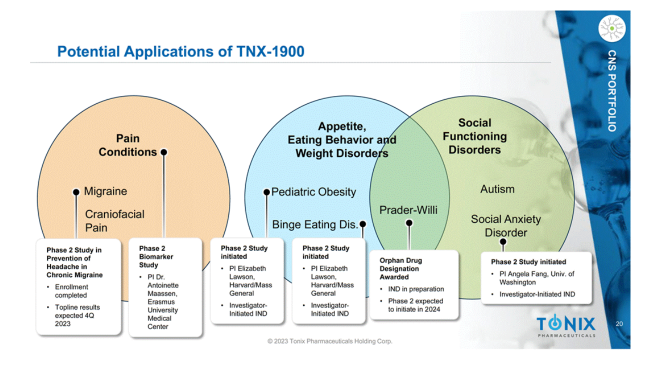
"A Phase 2 Study to Evaluate the Efficacy and Safety of TNX-102
SL in Patients With Multi-Site Pain Associated With Post-Acute
Sequelae of SARS-CoV-2 Infection (PREVAIL)

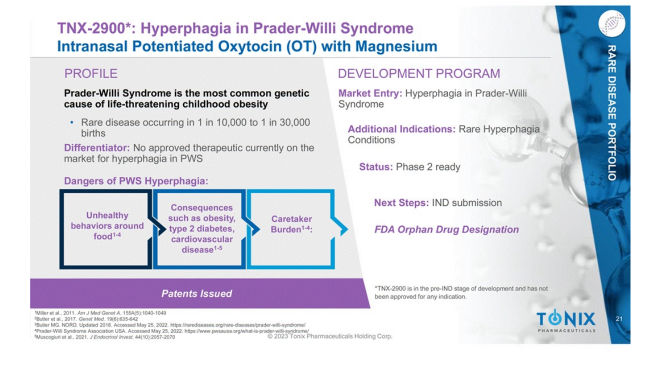
- 14 weeks -----

TONIX









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

PROFILE

- · A novel, oral, extended-release once-daily tablet
- Treatment effect of tianeptine sodium immediate release t.i.d. in depression is well-established
- Tianeptine restores neuroplasticity in animal models
- PPAR-β/δ and PPAR-y agonist1

Differentiators:

Relative to tianeptine IR available ex-US:

· Once daily dosing

Relative to traditional antidepressants:

- Unique mechanism of action beyond neurotransmitter modulation
- Tianeptine sodium IR has similar efficacy but less weight gain or sexual side effects than traditional antidepressants
- Tianeptine's side effects are described in labeling in countries in which it is marketed2

Patents Issued

DEVELOPMENT PROGRAM

Market Entry: Major Depressive Disorder (MDD)

Additional Indications: PTSD,

Neurocognitive Disorder From Corticosteroids, Alzheimer's Disease3

> Status: Phase 2 MDD study UPLIFT enrollment complete

Next Steps:

Topline results expected 4Q 2023

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

Sullivan G et al. Poster presentation at the American Society of Clinical Psychopharmacology, June 2023. https://bit.lyl42o3jnV/ /Summary of product characteristics (SmPC), European Medicines Agency, Stablon®, www.servier.ci/sites/default/files/spc-pil/SPC_Stablon_1.pdf accessed 7-16-23. /Garcia-Alberca et al., 2022. J Alzheimers Dis. 88(2):707-720

TNX-601 ER - Phase 2 UPLIFT* Study Design

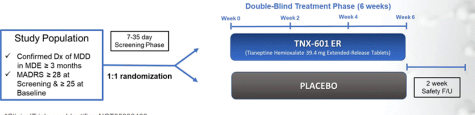
UPLIFTStudy

General study characteristics:

- Randomized, double-blind, placebo-controlled study in Major Depressive Disorder to evaluate monotherapy with TNX-601 ER versus placebo
- Parallel design with two arms treatment with tianeptine hemioxalate 39.4 mg or placebo
- U.S. sites only, completed enrollment of 116 patients

Primary Endpoint:

Mean change from baseline in the Montgomery-Åsberg Depression Rating Scale (MADRS) total score at Week 6



*ClinicalTrials.gov Identifier: NCT05686408

Abbreviations: Dx, diagnosis; ER, extended-release; F/U, follow-up; MDD, major depressive disorder; MDE, major depressive episode; N, number



Tianeptine is a Polyunsaturated Fatty Acid (PUFA) Analogue

PUFAs and PUFA-analogues

· Polar acidic "head" and hydrophobic fatty acid "tail"

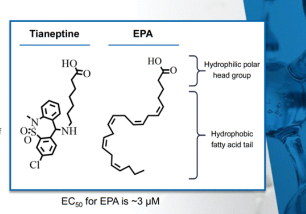
Pharmacological modulation of PUFA signaling has therapeutic potential in multiple pathologies

- · Act through the binding sites of endogenous FA metabolites on enzymes, transporters, and receptors
- · Several PUFA analogues have been developed as drugs, including the ethyl ester of eicosapentaenoic acid (EPA)1 which is branded as Vascepa® and Lovaza® (omega-3-acid ethyl esters)
- · Docosahexaenoic acid (DHA) is a primary structural component of

EPA and DHA have activity in treating MDD^{3,4} and Alzheimer's disease⁵

- · Activity shown in studies and meta-analyses, but insufficient randomized studies to be conclusive
- Pharmacology of EPA and DHA is not optimal⁵

¹EPA = eicosa-5Z, 8Z, 11Z, 14Z, 17Z-pentaenoic acid. ²Wikipedia: https://en.wikipedia.org/wiki/Docos ³Liao et al., 2019. *Transl Psychiatry*. 9(1):190



⁴Wani et al., 2015. *Integr Med Res.* 4(3):132-141 ⁵Heath RJ, and Wood TR. 2021. *Int J Mol Sci.* 2021 22(21):11826 © 2023 Tonix Pharmaceuticals Holding Corp.

TNX-601 ER - Racemic Tianeptine - Composed of Two Isomers

Racemic tianeptine:

- Approved in Europe and ex-US
- 1:1 mixture of 2 mirrorr-image isomers^{1,2}
- Weak µ-opioid receptor agonism²
 - Risk of abuse or diversion for euphoric effects3

	Racemic- Tianeptine	(S)- Tianeptine TNX-4300	(R)- Tianeptine
Activates PPAR-β/δ	+	+	-
Neuroplasticity	+	+	-
Novel Object Test ⁵	+	+	-
μ-Opioid Receptor	+	-	+
Forced Swim Test ⁶	+	-	+
Activates PPAR-y	+	+	+

(S)-Tianeptine: PPAR-β/δ agonist, no opiate liability4

New mechanism of action for treating depression



(R)-Tianeptine: opiate liability4

Weak µ-opioid receptor agonism4

(S)-tianeptine











Stablon. Summary of product characteristics. Les Laboratoires Servier Industrie; 2014.

*PubChem. Accessed November 10, 2022. https://pubchem.ncbi.nim.nih.gov/compound/Tianeptine

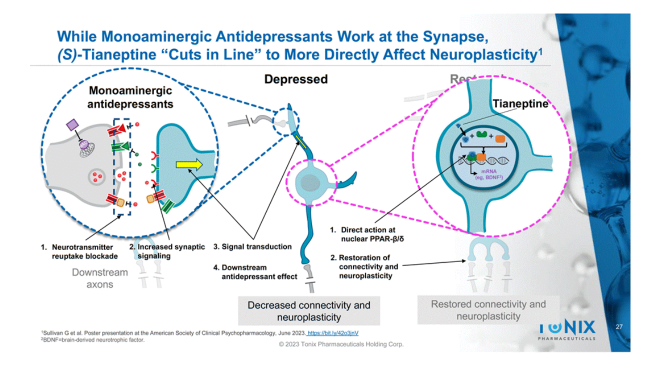
*Prug Enforcement Administration. May 2019. Accessed November 11, 2022. https://www.deadviversion.usdoi.gov/drug.chem.info/flaneptine.pdf

*Sallwan G et al. Poster presentation at the American Society of Clinical Psychopharmacology, June 2023. https://flat.lw/s2cSin/V

*Plat Novel Object Recognition Test

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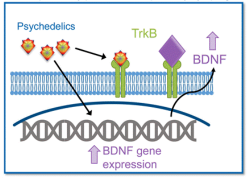
TNX-4300*: Depression, Alzheimer's & Parkinson's diseases Estianeptine (Single (S)-isomer of Tianeptine) **PROFILE DEVELOPMENT PROGRAM** · Single isomer, oral treatment Market Entry: Major Depressive Disorder (MDD) Proposed mechanism of action from lab studies indicates estianeptine is the active ingredient of TNX-601 ER1 Additional Indications: PTSD, PPAR-β/δ and PPAR-γ agonist Neurocognitive Disorder From Corticosteroids, Free of μ-opioid receptor activity Alzheimer's Disease² · Estianeptine restores neuroplasticity in tissue culture Status: Pre-clinical Differentiators: Next Steps: Expect IND can be Relative to racemic tianeptine IR or TNX-601 ER: supported by pre-clinical and clinical data from TNX-601 (racemic tianeptine) · Lack of opioid liability development Relative to traditional antidepressants: Unique mechanism of action – beyond neurotransmitter modulation Racemic tianeptine sodium IR has similar efficacy but fewer side effects than traditional antidepressants Patents Issued *TNX-4300 is in the pre-IND stage of development and has not been approved for any indication ¹Sullivan G et al. Poster presentation at the American Society of Clinical Psychopharmacology, June 2023. https://bit.ly/42o3jnV ²Garcia-Alberca et al., 2022. J Alzheimers Dis. 88(2):707-720

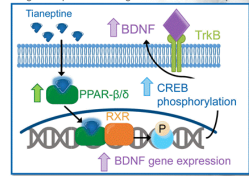


Tianeptine Shares a Neuroplasticity-Promoting Mechanism With Psychedelics

The neurotrophic growth factor BDNF plays a key role in the regulation of synaptic plasticity, and is diminished in populations suffering from anxiety and depression¹

- Psychedelics may promote neuroplasticity both by directly binding to BDNF receptor TrkB, and by increasing BDNF gene expression^{1,2}
- Tianeptine may promote neuroplasticity by upregulating BDNF gene expression through activation of PPAR-β/δ^{3,4}





BDNF=brain-derived neurotrophic factor: CREB=cAMP response elem 'de Vos CMH, et al. Front Psychiatry. 2021;12:724060 *Nolliner R, et al. Nat Neurosci. 2023;26(6):1032-1041 '31 MJ, et al. Int J Neuropsychopharmacol. 2015;19(1):pyv083 '85eo MK, et al. Psychopharmacology (Berl). 2016;233(13):2617-2627

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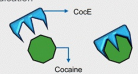
TNX-1300*: Cocaine Intoxication Cocaine Esterase (CocE)

PROFILE

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

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

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





Patents Issued

DEVELOPMENT PROGRAM

Market Entry: Cocaine Intoxication

Status: Mid-Phase 2

Next Steps: Initiate new Phase 2 trial 3Q

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

FDA Breakthrough Therapy Designation

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

*TNX-1300 has not been approved for any indication

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







Next Generation α -CD40 Ligand (CD40L) Antibody

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

Differentiators: Expected to deliver efficacy without compromising safety

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

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

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

TNX-1500 has not been approved for any indication. Patents filed

Prevention of Allograft Rejection

Status: Phase 1 ready - IND cleared

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

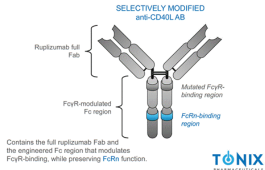
Next Steps: Initiate Phase 1 study 3Q 2023

Autoimmune Diseases

Status: Potential future indications include:

Sjögren's Syndrome, Systemic Lupus Erythematosus

These indications require large studies, but represent large target markets



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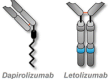
Third-Generation α-CD40L **Engineered to Decrease Risk of Thrombosis**

First-generation anti-CD40L mAbs Ruplizumab

Constant fragment (Fc) domain interacted with FcvRIIA (CD32A). which suggested a mechanism for the increased risk of thrombosis. 1,2

Second-generation anti-CD40L proteins







Second-generation anti-CD40L proteins exhibited dramatically reduced binding to FcyRIIA3-6 but had other issues, including decreased efficacy, shortened half-life, or engendering of anti-drug antibodies (ADAs).7-9

Third-generation anti-CD40L mAbs*



TNX-1500

TNX-1500 is engineered to target CD40L therapeutically while reducing FcγRIIA binding and thereby lowering the potential for thrombosis.1-5

*Sanofi's frexalimab (formerly SAR441344) and Eledon's tegoprubart (formerly AT-1501) also are Fc modified

-mward et al., 2003. Circ Res. 92(9):1041-1048
-Robles-Carrillo et al., 2010. J Immunol. 185(3):1577-1583
-Shock et al., 2015. Arthritis Res Ther. 17(1):234
-Kke et al., 2014. J Immunol. 19(1):1583-1594
-Kennell et al., 2004. Int Immunol. 16(11):1583-1594
-Kannell et al., 2019. Sci Trans Med 11(489):eaan584
-ClinicarTrials gov identifier. NCT02273960. Updated July 16, 2019. Accessed June 1, 2021. https://clinicaltrials.gov/ct2/show/results/NCT02273960?view=results
-Waters. 2018. Biocentury.
-Company data

Aglycosyl

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Other anti-CD40L Monoclonal Antibodies in Development



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

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



Horizon (being acquired by Amgen) - Sjögren's Syndrome (SjS)

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

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

- Phase 2 Trial Currently Enrolling in SjS (NCT04572841) and SLE (NCT05039840)
- Active Phase 2 Trial in Relapsing MS (NCT04879628)
- · Frexalimab, f.k.a.SAR441344 (Fc-modified)



Eledon - Amyotrophic Lateral Sclerosis (ALS) and Kidney Transplant

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



Lundbeck and AprilBio - Neurology

- Phase 1 Trial Currently Enrolling in Healthy Adults (NCT05136053)
- APB-A1 or Lu AG22515 (HAS fusion protein)

https://www.ucb.com/our-science/pipeline
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IMMUNOLOGY PORTFOLIO

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

Potential New Cancer Treatment

- TNX-1700 (rTFF2-HSA) has effects on cancer by altering the tumor micro-environment
- Mechanism of action: suppresses myeloid-derived suppressor cells and activates anti-cancer CD8+ T cells
- Potential synergy with anti-PD1 or anti-PD-L1 monoclonal antibodies (mAbs)

Preclinical Evidence for Inhibiting Growth of Cancer Cells

- In an MC38 mouse model of colorectal cancer, mTNX-1700 (murine TNX-1700) alone inhibited tumor growth by 50%, and combination therapy with anti-PD1 inhibited tumor growth by 87%¹
- In an advanced mouse model of gastric cancer, mTNX-1700 combination therapy with anti-PD1 inhibited tumor growth by 78%²
- Mechanistically, the combination therapy reduced intratumoral MDSCs, profoundly increased tumor-infiltrating CD8+ T cells, and significantly reduced spontaneous metastasis²

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

Status: Preclinical

Next Steps: Animal studies ongoing

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

Licensed from Columbia University

 Developing in partnership under sponsored research agreement

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

Patents Filed

Daugherty et al., AACR Poster. 2023. MDSC-targeted TFF2-MSA suppresses tumor growth and increases survival in anti-PD-1 treated MC38 and CT26.wt murine colorectal cancer models. https://bit.hu45xbGK9

https://bit.hids.thids.bt.GK9
**Olan et al., AACR Poster. 2023. MDSC-targeted TFF2-MSA synergizes with PD-1 blockade therapy in diffuse-type gastric cancer. https://bit.hy/lacQsku
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TNX-801*



Recombinant Pox Vaccine (RPV)
Platform Using Live Virus Technology

Differentiators:

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

TNX-801 is in the pre-IND stage of development and has not been approved for any indication. Patents file

Noyce et al., 2018. PLoS One. 13(1):e01884

Mpox and Smallpox Vaccine

Status: Preclinical

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

Next Steps: File IND



Vaccine for Future Emerging Infectious Diseases

Example: TNX-1850 for COVID-19

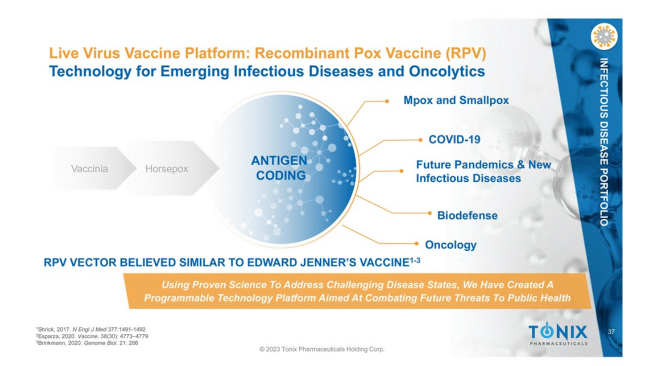
Status: Model System

TNX-801* scHPXV (Horsepox) 212,811 bp





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Internal Development & Manufacturing Capabilities

R&D Center (RDC) - Frederick, MD

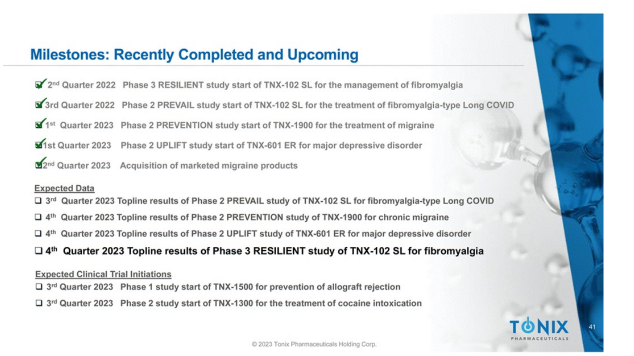
- · Functions:
 - Research advancing CNS and immunology drugs
 - Accelerated development of vaccines and antiviral drugs against COVID-19, its variants and other infectious diseases
- Description: ~48,000 square feet, BSL-2 with some areas designated BSL-3
- · Status: Operational

Advanced Development Center (ADC) - North Dartmouth, MA

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









Zembrace® IMPORTANT SAFETY INFORMATION (1 of 2)

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

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

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

Do not use Zembrace if you have:

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

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

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



Zembrace® IMPORTANT SAFETY INFORMATION (2 of 2)

Zembrace may cause serious side effects including:

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

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

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

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

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

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

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

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Tosymra® IMPORTANT SAFETY INFORMATION (1 of 2)

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

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

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

Do not use Tosymra if you have:

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

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

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Tosymra® IMPORTANT SAFETY INFORMATION (2 of 2)

Tosymra may cause serious side effects including:

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

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

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

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

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

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

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

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

