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 14, 2017

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

Copy of correspondence to:

Marc J. Ross, Esq.
James M. Turner, Esq.
Sichenzia Ross Ference Kesner LLP
61 Broadway
New York, New York 10006
Tel: (212) 930-9700 Fax: (212) 930-9725

Check the appropriate	e box below	if the Form	8-K fi	ling is	intended	to si	imultaneously	satisfy	the	filing	obligation	of the	e registrant	under
any of the following n	rovisions (s	ee General Ir	structi	on A.2	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))

Item 8.01 Other Events.

On March 14, 2017, Tonix Pharmaceuticals Holding Corp. (the "Company") issued a press release announcing that the United States Patent and Trademark Office has issued a Notice of Allowance for U.S. Patent Application 14/214,433, "Eutectic Formulations of Cyclobenzaprine Hydrochloride and Amitriptyline Hydrochloride," covering the proprietary sublingual formulation of TNX-102 SL.

A copy of the press release that discusses this matter is filed as Exhibit 99.01 to, and incorporated by reference in, this report. The information in this Current Report is being furnished and shall not be deemed "filed" for the purposes of Section 18 of the Securities Exchange Act of 1934 or otherwise subject to the liabilities of that Section. The information in this Current Report shall not be incorporated by reference into any registration statement or other document pursuant to the Securities Act of 1933, except as shall be expressly set forth by specific reference in any such filing.

Item 9.01 Financial Statements and Exhibits.

(d) Exhibits.

99.01 Press release, dated March 14, 2017, issued by Tonix Pharmaceuticals Holding Corp.*

^{*} Furnished herewith.

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 14, 2017 By: /s/ SETH LEDERMAN

Seth Lederman

Chief Executive Officer



Tonix Pharmaceuticals Receives Notice of Allowance for New U.S. Patent Covering Composition and Manufacture of TNX-102 SL

Patent Will Provide Intellectual Property Protection until 2034 to TNX-102 SL, an FDA-Designated Breakthrough Therapy in Phase 3
Development for Posttraumatic Stress Disorder (PTSD)

NEW YORK, Mar. 14, 2017 (GLOBE NEWSWIRE) – <u>Tonix Pharmaceuticals Holding Corp.</u> (Nasdaq:TNXP) (Tonix), a company that is developing innovative pharmaceutical products to address public health challenges, announced today that the U.S. Patent and Trademark Office has issued a Notice of Allowance for U.S. Patent Application 14/214,433, "Eutectic Formulations of Cyclobenzaprine Hydrochloride and Amitriptyline Hydrochloride," covering the proprietary sublingual formulation of TNX-102 SL*. A Notice of Allowance signifies that Tonix will be entitled to receive patent protection until 2034 in the U.S. for the allowed claims when the patent is issued. Tonix expects the patent to be issued within two months.

The allowed claims protect the pharmaceutical composition and the method of manufacturing of TNX-102 SL. The TNX-102 SL sublingual formulation is based on a eutectic between cyclobenzaprine HCl and mannitol, which protects the acidic hydrochloride salt of cyclobenzaprine from molecular interactions with the basic excipient, potassium phosphate dibasic, which is added to enhance transmucosal absorption. Transmucosal absorption of cyclobenzaprine increases the rate of absorption into the blood stream and bypasses first pass liver metabolism. TNX-102 SL is distinct from orally ingested forms of cyclobenzaprine, which are available as generic immediate-release tablets and branded extended-release capsules (AMRIX®), and are approved for short-term use (2-3 weeks) for relief of muscle spasm associated with acute, painful musculoskeletal conditions. Since TNX-102 SL has a different route of administration and different pharmacokinetic profile from orally ingested cyclobenzaprine and is intended for a new indication (PTSD), pharmacists will not be able to substitute orally ingested forms of cyclobenzaprine for TNX-102 SL.

Tonix has filed trademarks to describe the protective eutectic, ProtecticTM, and the formulation that utilizes interactions at the angstrom scale, Angstro-TechnologyTM. The ProtecticTM protective eutectic and Angstro-TechnologyTM formulation enable TNX-102 SL to be a sublingual tablet of cyclobenzaprine HCl, while ensuring that the tablets have robust manufacturability and pharmaceutical stability.

Seth Lederman, M.D., president and chief executive officer of Tonix, stated, "This notice of allowance strengthens the value of our pipeline with near-term intellectual property protection and highlights Tonix's innovation in proprietary pharmaceutical development. Most importantly, the ProtecticTM protective eutectic and Angstro-TechnologyTM formulation are essential elements of the proprietary TNX-102 SL composition."

*TNX-102 SL (cyclobenzaprine HCl sublingual tablets) is an investigational new drug and has not been approved for any indication.

About Tonix Pharmaceuticals Holding Corp.

Tonix is developing innovative pharmaceutical products to address public health challenges. TNX-102 SL is in Phase 3 development and has been granted Breakthrough Therapy designation by the U.S. Food and Drug Administration (FDA) for the treatment of PTSD. PTSD is a serious condition characterized by chronic disability, inadequate treatment options especially for military-related PTSD, and an overall high utilization of healthcare services that contributes to significant economic burdens. Other development efforts include TNX-601 (tianeptine oxalate), a clinical candidate at Pre-IND (Investigational New Drug) application stage, designed for daytime use for the treatment of PTSD, and TNX-801, a potential smallpox-preventing vaccine based on a live synthetic version of horsepox virus (HPXV). HPXV has protective vaccine activity in mice, using a model of lethal vaccinia infection. Vaccine manufacturing activities have been initiated to support further nonclinical testing of TNX-801.

This press release and further information about Tonix can be found at www.tonixpharma.com.

Forward Looking Statements

Certain statements in this press release are forward-looking within the meaning of 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," "expect," 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 FDA clearances or approvals and noncompliance with FDA regulations. As with any pharmaceutical under development, there are significant risks in the development, regulatory approval and commercialization of new products. Tonix does not undertake an obligation to update or revise any forward-looking statement. Investors should read the risk factors set forth in the Annual Report on Form 10-K for the year ended December 31, 2015, as filed with the Securities and Exchange Commission (the "SEC") on March 3, 2016, and future periodic reports filed with the SEC on or after the date hereof. All of Tonix's forward-looking statements are expressly qualified by all such risk factors and other cautionary statements. The information set forth herein speaks only as of the date hereof.

Contacts

Jessica Smiley Investor Relations investor.relations@tonixpharma.com (212) 980-9155 x185

Edison Advisors (investors) Tirth Patel tpatel@edisongroup.com (646) 653-7035

Russo Partners (media) Rich Allan <u>rich.allan@russopartnersllc.com</u> (646) 942-5588