
**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): **May 29, 2025**

ACURA PHARMACEUTICALS, INC.
(Exact Name of Registrant as specified in its Charter)

New York
(State or other jurisdiction of
incorporation or organization)

1-10113
(Commission File Number)

11-0853640
(I.R.S. Employer
Identification Number)

616 N. North Court, Suite 120
Palatine, Illinois 60067
(Address of principal executive offices) (Zip code)

(847) 705-7709
(Registrant's telephone number, including area code)

Securities registered pursuant to Section 12(b) of the Act:

<u>Title of Each Class</u>	<u>Trading Symbol(s)</u>	<u>Name of Each Exchange on Which Registered</u>
Common Stock, \$0.01 par value per share	ACUR	OTC Market – OTC Expert Market

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:

- 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 (17CFR240.14d-2(b))
- Pre-commencement communications pursuant to Rule 13e-4(c) under the Exchange Act (17CFR 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

If an emerging growth company, indicate by check mark if the registrant has elected not to use the extended transition period for complying with any new or revised financial accounting standards provided pursuant to Section 13(a) of the Exchange Act.

Item 1.01 - Entry into a Material Definitive Agreement.

On both May 2, 2025 and May 22, 2025, we received \$100,000 loans from Abuse Deterrent Pharma, LLC (“AD Pharma”). These loans combined with previous loans made to the Company and combined with the \$2,319,279 under the November 10, 2022 Amended Consolidated and Restated Secured Promissory Note, now has a principal balance of \$7,994,279 with accrued interest of approximately \$683,000 as of May 29, 2025 and bears interest at 5.25% (“Note”). The Events of default under the Note include, among other items, bankruptcy events, failure to pay interest and principal when due and such failure continues for 5 days, and if Acura is generally not, or is unable to, or admits in writing its inability to, pay its debts as those debts become due. If any amount payable hereunder is not paid when due (without regard to any applicable grace periods), whether at stated maturity, by acceleration, or otherwise, including upon an event of default, such overdue amount shall bear interest at the rate per annum of 7.5% from the date of such non-payment until such amount is paid in full.

The funding provided by AD Pharma will be used to meet day-to-day operation activity. There can be no assurance we will be successful in receiving additional financing. In the absence of the receipt of additional financing by mid-June 2025, we will be required to scale back our operations, including the furlough and lay-off of employees, or to terminate operations and/or seek protection under applicable bankruptcy laws. This could result in a complete loss of shareholder value in the Company. Even assuming we are successful in securing additional sources of financing to fund continued operations, there can be no assurance that the proceeds of such financing will be sufficient to fund operations until such time, if at all, that we generate sufficient revenue from our products and product candidates to sustain and grow our operation.

The inclusion of a description of the Note under Item 1.01 of this Current Report on Form 8-K shall not be deemed an acknowledgement that the Note is a material agreement not made, or deemed not to be made, in the ordinary course of our business.

Item 1.01 - Entry into a Material Definitive Agreement.

On May 29, 2025, Acura Pharmaceuticals, Inc. (“we” “Acura” or the “Company”), received the executed agreement to further amend the June 28, 2019 License, Development and Commercialization Agreement (“Amended Agreement”) with AD Pharma, for the development of LTX-03 (hydrocodone bitartrate with acetaminophen) immediate-release tablets utilizing Acura’s patented LIMITx technology which addresses the consequences of excess oral administration of opioid tablets, the most prevalent route of opioid overdose and abuse.

This amendment #12 has an effective date of May 29, 2025 and extends the FDA’s acceptance date of a New Drug Application (“NDA”) for LTX-03 from May 31, 2025 to December 31, 2025 (“NDA Acceptance Date”) (“Amended Agreement”).

AD Pharma may terminate the Amended Agreement at any time. Additionally, if the NDA for LTX-03 is not accepted by the FDA by the NDA Acceptance Date, AD Pharma may terminate the Amended Agreement and take ownership of the intellectual property rights of LTX-03 from the Company. Should AD Pharma choose not to exercise this option to terminate the Amended Agreement and the NDA for LTX-03 is subsequently accepted by the FDA, such option to terminate the Amended Agreement expires.

The inclusion of a description of the Amended Agreement with AD Pharma under Item 1.01 of this Current Report on Form 8-K shall not be deemed an acknowledgement that the Amended Agreement is a material agreement not made, or deemed not to be made, in the ordinary course of our business.

Item 1.01 - Entry into a Material Definitive Agreement.

On May 29, 2025, we received the executed agreement to further amend the November 10, 2022 Amended, Consolidated and Restated Secured Promissory Note (the “Amended Note”) with AD Pharma. This amendment #6 has an effective date of May 29, 2025 and changes the maturity date of the Amended Note from May 31, 2025 to December 31, 2025, at which time all principal and interest is due.

The inclusion of a description of the Amended Note under Item 1.01 of this Current Report on Form 8-K shall not be deemed an acknowledgement that the Amended Note is a material agreement not made, or deemed not to be made, in the ordinary course of our business.

Item 1.01 – Entry into a Material Definitive Agreement.

We previously announced on June 28, 2019, we entered into a promissory note with John Schutte that consolidated existing promissory notes into a single note for \$6.0 million bearing interest at 7.5% (“\$6.0 Million Note”). At that time, we also granted to Mr. Schutte conversion rights of the \$6.0 Million Note into Acura common stock at \$0.16 per share, issued to him a warrant to purchase 10.0 million shares of the Company’s common stock at a price of \$0.01 per share having an expiration date of June 28, 2024 (“Warrant”) and granted a security interest in all our assets (“Security Agreement”). With our consent, effective on June 28, 2019, Mr. Schutte assigned and transferred to AD Pharma all of his right, title and interest in this \$6.0 Million Loan, Security Agreement and Warrant. We previously announced on June 9, 2021, we received notice of conversion from AD Pharma for the \$6.0 Million Note and approximately \$877 thousand of accrued but unpaid interest on the \$6.0 Million Note. The principal and interest on the \$6.0 Million Note were converted into 42,984,375 shares of the Company’s common stock.

Effective May 29, 2025, in consideration of further amending the November 10, 2022 Amended, Consolidated and Restated Secured Promissory Note (the “Note”) with AD Pharma which changed the maturity date of the Note from May 31, 2025 to December 31, 2025, we amended the Warrant to change the expiration date of the Warrant from May 31, 2025 to December 31, 2025.

The inclusion of a description of the Warrant under Item 1.01 of this Current Report on Form 8-K shall not be deemed an acknowledgement that the Note is a material agreement not made, or deemed not to be made, in the ordinary course of our business.

At May 29, 2025, AD Pharma directly owns approximately 65% of the outstanding common stock of the Company. The ownership percentage of the Company held by AD Pharma does not include their warrant to purchase 10.0 million shares of common stock of the Company. AD Pharma is an entity controlled by Mr. Schutte, of which Mr. Schutte is the managing partner and investor. At May 29, 2025, Mr. Schutte directly owns approximately 13% of the outstanding common stock of the Company.

Item 2.01 – Completion of Acquisition or Disposition of Assets

The contents of all Items 1.01 are incorporated herein by reference.

Item 2.03 - Creation of a Direct Financial Obligation or an Obligation under an Off-Balance Sheet Arrangement of a Registrant

The contents of all Items 1.01 are incorporated herein by reference.

Acura Forward-Looking Statements

Statements in this Current Report constitute forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995 and these forward-looking statements are made in reliance on the safe harbor provisions of Section 27A of the Securities Act of 1933, as amended, and Section 21E of the Securities Exchange Act of 1934, as amended. Such forward-looking statements involve known and unknown risks, uncertainties and other factors which may cause our actual results, performance or achievements to be materially different from any future results, performance, or achievements expressed or implied by such forward-looking statements.

Forward-looking statements may include, but are not limited to:

- whether the FDA will agree with or accept the results of our studies for our product candidates;
- the ability to fulfill the FDA requirements for approving our product candidates for commercial manufacturing and distribution in the United States, including, without limitation, the adequacy of the results of the laboratory and clinical studies completed to date, the results of laboratory and clinical studies we may complete in the future to support FDA approval of our product candidates and the sufficiency of our development process to meet over-the-counter (“OTC”) Monograph standards, as applicable;
- whether we can successfully submit a New Drug Application for LTX-03, request a priority review and whether such filings and requests will be accepted by the FDA;
- our ability to obtain funding from Abuse Deterrent Pharma, LLC or other parties for our continuing operations, including the development of our products utilizing our LIMITx™ and Impede® technologies;
- whether we can renegotiate the date by which we are required to obtain FDA acceptance, currently December 31, 2025, for an NDA for LTX-03 by our agreement with Abuse Deterrent Pharma, LLC on which we depend to finance operations;
- whether we can renegotiate the date by which we are required to pay off the secured promissory note and accrued interest to Abuse Deterrent Pharma, LLC, currently December 31, 2025;
- whether our licensing partners will develop any additional products and utilize Acura for such development;
- the expected results of clinical studies relating to LTX-03, a LIMITx hydrocodone bitartrate and acetaminophen combination product, or any successor product candidate, the date by which such studies will be complete and the results will be available and whether LTX-03 will ultimately receive FDA approval;
- our business could be adversely affected by health epidemics in regions where third parties for which we rely, as in CROs or CMOs, have concentrations of clinical trial sites or other business operations, and could cause significant disruption in the operations of third-party manufacturers and CROs upon whom we rely;

- whether LIMITx will retard the release of opioid active ingredients as dose levels increase;
- whether the extent to which products formulated with the LIMITx Technology reduce respiratory depression will be determined sufficient by the FDA to support approval or labelling describing safety features;
- our and our licensee's ability to successfully launch and commercialize our products and technologies;
- the results and timing of our development of our LIMITx Technology, including, but not limited to, the submission of a NDA and/or FDA filing acceptance;
- our or our licensees' ability to obtain necessary regulatory approvals and commercialize products utilizing our technologies;
- the market acceptance of, timing of commercial launch and competitive environment for any of our products;
- expectations regarding potential market share for our products;
- our ability to develop and enter into additional license agreements for our product candidates using our technologies;
- our exposure to product liability and other lawsuits in connection with the commercialization of our products;
- the increasing cost of insurance and the availability of product liability insurance coverage;
- the ability to avoid infringement of patents, trademarks and other proprietary rights of third parties;
- the ability of our patents to protect our products from generic competition and our ability to protect and enforce our patent rights in any paragraph IV patent infringement litigation;
- the adequacy of the development program for our product candidates, including whether additional clinical studies will be required to support FDA approval of our product candidates;
- changes in regulatory requirements;
- adverse safety findings relating to our commercialized products or product candidates in development;
- whether the FDA will agree with our analysis of our clinical and laboratory studies;
- whether or when we are able to obtain FDA approval of labeling for our product candidates for the proposed indications and whether we will be able to promote the features of our technologies; and
- whether our product candidates will ultimately perform as intended in commercial settings.

In some cases, you can identify forward-looking statements by terms such as "aim", "anticipate", "believe", "could", "design", "estimate", "expect", "forecast", "goal", "guidance", "imply", "indicate", "intend", "may", "objective", "opportunity", "outlook", "plan", "position", "potential", "predict", "project", "prospective", "pursue", "seek", "should", "strategy", "target", "would", "will", and other words of similar meaning, expressions, derivations of such words and the use of future dates intended to identify forward-looking statements. These statements reflect our current views with respect to future events and are based on assumptions and subject to risks and uncertainties. Given these uncertainties, you should not place undue reliance on these forward-looking statements. We discuss many of these risks in greater detail in Acura's Annual Report on Form 10-K and Quarterly Reports on Form 10-Q filed with the U.S. Securities and Exchange Commission ("SEC") and in other filings Acura makes with the SEC from time to time. Investors and potential investors are urged not to place undue reliance on forward-looking statements in this communication, which speak only as of this date of the Current Report and are based on the Company's current beliefs, assumptions, and expectations. While Acura may elect to update these forward-looking statements at some point in the future, it specifically disclaims any obligation to update or revise any forward-looking statements contained in this Current Report whether as a result of new information or future events, except as may be required by applicable law.

Item 9.01 - Financial Statements and Exhibits

<u>Exhibit Number</u>	<u>Description</u>
99.1	Amended Loan Schedule dated May 22, 2025 to the November 10, 2022 Amended, Consolidated and Restated Secured Promissory Note with Abuse Deterrent Pharma, LLC
99.2	Amendment #12 dated May 29, 2025 to the License, Development and Commercialization Agreement with Abuse Deterrent Pharma, LLC
99.3	Amendment #6 dated May 29, 2025 to the November 10, 2022 Amended, Consolidated and Restated Secured Promissory Note with Abuse Deterrent Pharma, LLC
99.4	Amendment #4 dated May 29, 2025 to the June 28, 2019 Common Stock Purchase Warrant with Abuse Deterrent Pharma, LLC
104	Cover Page Interactive Data File (embedded within Inline XBRL document)

SIGNATURES

Pursuant to the requirements 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.

ACURA PHARMACEUTICALS, INC.

By: /s/ Robert A. Seiser
Robert A. Seiser
Senior Vice President & Chief Financial Officer

Date: May 30, 2025

**Amended Loan Schedule to Secured Promissory Note dated November 10, 2022
between Acura Pharmaceuticals, Inc. and Abuse Deterrent Pharma, LLC**

	<u>Date</u>	<u>Principal</u>	<u>Aggregated Principal</u>
Original Secured Promissory Note	11/10/2022	\$2,319,279	\$2,319,279
Additional Loans to be included:			
Loans #1 dated 12/22/2022 through Loan #26 dated 12/12/2024		\$ 4,625,000	\$ 6,944,279
Loan #27	1/13/2025	\$ 125,000	\$ 7,069,279
Loan #28	1/23/2025	\$ 100,000	\$ 7,169,279
Loan #29	2/03/2025	\$ 125,000	\$ 7,294,279
Loan #30	2/14/2025	\$ 100,000	\$ 7,394,279
Loan #31	2/28/2025	\$ 100,000	\$ 7,494,279
Loan #32	3/17/2025	\$ 100,000	\$ 7,594,279
Loan #33	4/04/2025	\$ 100,000	\$ 7,694,279
Loan #34	4/18/2025	\$ 100,000	\$ 7,794,279
Loan #35	5/02/2025	\$ 100,000	\$ 7,894,279
Loan #36	5/22/2025	\$ 100,000	\$ 7,994,279

ACURA PHARMACEUTICALS, INC.

By: /s/ Robert A. Seiser

Robert A. Seiser
Senior Vice President & CFO
Date: May 22, 2025

**AMENDMENT #12
TO LICENSE, DEVELOPMENT AND COMMERCIALIZATION AGREEMENT**

This AMENDMENT #12 (this "**Amendment**") TO LICENSE, DEVELOPMENT AND COMMERCIALIZATION AGREEMENT (the "**Agreement**") dated June 28, 2019 between Acura Pharmaceuticals, Inc. ("**Acura**"), a New York corporation, having a place of business at 616 N. North Court, Suite 120, Palatine, IL 60067, and Abuse Deterrent Pharma, LLC ("**AD Pharma**"), a Kentucky limited liability company, having a place of business at 2604 River Green Circle, Louisville, Kentucky 40206, is made as of May 29, 2025.

RECITALS

WHEREAS, Acura and AD Pharma have entered into certain Amendments to the Agreement (the "**Prior Amendments**") as follows:

Amendment # 1 on October 16, 2020, Amendment #2 on June 17, 2021,
Amendment #3 on February 28, 2022,
Amendment #4 on November 10, 2022,
Amendment #5 on December 8, 2022,
Amendment #6 on June 15, 2023,
Amendment #7 on November 13, 2023,
Amendment #8 on March 15, 2024,
Amendment #9 on June 14, 2024,
Amendment #10 on October 28, 2024
Amendment #11 on February 28, 2025, **and**

WHEREAS, the Parties desire to amend the Agreement to provide for an extension to the LIMITx™ Regulatory Submission Timeline.

NOW THEREFORE, in consideration of the mutual covenants and promises contained in the Agreement, as amended, and this Amendment and other good and valuable consideration, the receipt and sufficiency of which are hereby acknowledged, Acura and AD Pharma agree as follows:

**ARTICLE 1
AMENDMENTS TO AGREEMENT**

1.1 Item 3 of Schedule 1 "LIMITx™ Regulatory Application Submission Timeline" is hereby amended and replaced in its entirety as follows:

3. By December 31, 2025 Acura must gain filing acceptance by the FDA of a Regulatory Approval Application for the Product.

**ARTICLE 2
MISCELLANEOUS**

2.1 Governing Law. This Amendment shall be governed by the laws of the State of New York without regard to its conflict of laws rules or principles.

2.2 Amendments. Except as expressly amended by this Amendment #12 and the "Prior Amendments", the Agreement shall remain unmodified and in full force and effect.

2.3 Entire Agreement. The Agreement (including the Schedules attached thereto), as amended by the "Prior Amendments" and this Amendment, constitutes the entire agreement of the Parties with respect to the subject matter hereof and supersede all prior understandings and writings between the Parties relating thereto.

2.4 Interpretation. Any capitalized terms used in this Amendment and not otherwise defined herein shall have the meaning provided in the Agreement.

2.5 Counterparts. This Amendment may be executed manually or electronically by the Parties, in any number of counterparts, each of which shall be considered one and the same amendment and shall become effective when a counterpart hereof shall have been signed by each of the Parties and delivered to the other Party.

IN WITNESS WHEREOF, the Parties hereto have caused this Amendment to be executed in their names by their properly and duly authorized officers or representatives as of the date first written above.

ACURA PHARMACEUTICALS, INC.

By: /s/ Robert A. Seiser
Name: Robert Seiser
Title: SVP and CFO

ABUSE DETERRENT PHARMA, LLC

By: /s/ John Schutte
Name: John Schutte
Title: Managing Partner

**AMENDMENT #6
AMENDED, CONSOLIDATED AND RESTATED
SECURED PROMISSORY NOTE**

This AMENDMENT #6 (this “**Amendment**”) TO AMENDED, CONSOLIDATED AND RESTATED SECURED PROMISSORY NOTE (the “**Note**”) dated November 10, 2022 between Acura Pharmaceuticals, Inc. (“**Acura**”), a New York corporation, having a place of business at 616 N. North Court, Suite 120, Palatine, IL 60067, and Abuse Deterrent Pharma, LLC (“**AD Pharma**”), a Kentucky limited liability company, having a place of business at 2604 River Green Circle, Louisville, Kentucky 40206, is made as of May 29, 2025.

RECITALS

WHEREAS, the Parties desire to amend the Note to change the Maturity Date of this Note.

NOW THEREFORE, in consideration of the mutual covenants and promises contained in the Note, as amended, and this Amendment and other good and valuable consideration, the receipt and sufficiency of which are hereby acknowledged, Acura and AD Pharma agree as follows:

**ARTICLE 1
AMENDMENT TO AGREEMENT**

1.1 Item 2.1 “Maturity Date” is hereby amended and replaced in its entirety as follows:

The Company agrees to pay the principal sum of this Note and interest on the unpaid principal sum of this Note on December 31, 2025 (the “**Maturity Date**”). Time shall be of the essence with respect to all of the Company’s obligations under this Note.

2.1 Amendments. Except as expressly amended by this Amendment #6 and the “Prior Amendments”, the Note shall remain unmodified and in full force and effect.

IN WITNESS WHEREOF, the Parties hereto have caused this Amendment to be executed in their names by their properly and duly authorized officers or representatives as of the date first written above.

ACURA PHARMACEUTICALS, INC.

By: /s/ Robert A. Seiser

Name: Robert Seiser

Title: SVP and CFO

ABUSE DETERRENT PHARMA, LLC

By: /s/ John Schutte

Name: John Schutte Title: Managing Partner

AMENDMENT #4 COMMON STOCK PURCHASE WARRANT

This AMENDMENT #4 (this “**Amendment**”) TO COMMON STOCK PURCHASE WARRANT (the “**Warrant**”) dated June 28, 2019 between Acura Pharmaceuticals, Inc. (“**Acura**”), a New York corporation, having a place of business at 616 N. North Court, Suite 120, Palatine, IL 60067, and Abuse Deterrent Pharma, LLC (“**AD Pharma**”), a Kentucky limited liability company, having a place of business at 2604 River Green Circle, Louisville, Kentucky 40206, is made as of May 29, 2025.

RECITALS

WHEREAS, the Parties desire to amend the Warrant to change the Expiration Date of this Warrant.

NOW THEREFORE, in consideration of the mutual covenants and promises contained in the Warrant, as amended, and this Amendment and other good and valuable consideration, the receipt and sufficiency of which are hereby acknowledged, Acura and AD Pharma agree to change Expiration Date of this Warrant to December 31, 2025.

Amendments. Except as expressly amended by this Amendment #4, the Warrant shall remain unmodified and in full force and effect.

IN WITNESS WHEREOF, the Parties hereto have caused this Amendment to be executed in their names by their properly and duly authorized officers or representatives as of the date first written above.

ACURA PHARMACEUTICALS, INC.

By: /s/ Robert A. Seiser
Name: Robert Seiser
Title: SVP and CFO

ABUSE DETERRENT PHARMA, LLC

By: /s/ John Schutte
Name: John Schutte
Title: Managing Partner
